Review

Select Polyphenol-Rich Berry Consumption to Defer or Deter Diabetes and Diabetes-Related Complications

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Abstract: Berries are considered “promising functional fruits” due to their distinct and ubiquitous therapeutic contents of anthocyanins, proanthocyanidins, phenolic acids, flavonoids, flavanols, alkaloids, polysaccharides, hydroxycinnamic, ellagic acid derivatives, and organic acids. These polyphenols are part of berries and the human diet, and evidence suggests that their intake is associated with a reduced risk or the reversal of metabolic pathophysiology related to diabetes, obesity, oxidative stress, inflammation, and hypertension. This work reviewed and summarized both clinical and non-clinical findings that the consumption of berries, berry extracts, purified compounds, juices, jams, jellies, and other berry byproducts aided in the prevention and or otherwise management of type 2 diabetes mellitus (T2DM) and related complications. The integration of berries and berries-derived byproducts into high-carbohydrate (HCD) and high-fat (HFD) diets, also reversed/reduced the HCD/HFD-induced alterations in glucose metabolism-related pathways, and markers of oxidative stress, inflammation, and lipid oxidation in healthy/obese/diabetic subjects. The berry polyphenols also modulate the intestinal microbiota ecology by opposing the diabetic and obesity rendered symbolic reduction of Bacteroidetes/Firmicutes ratio, intestinal mucosal barrier dysfunction-restoring bacteria, short-chain fatty acids, and organic acid producing microflora. All studies proposed a number of potential mechanisms of action of respective berry bioactive compounds, although further mechanistic and molecular studies are warranted. The metabolic profiling of each berry is also included to provide up-to-date information regarding the potential anti-oxidative/antidiabetic constituents of each berry.

Keywords: berries; metabolic syndrome; precision nutrition; hyperglycemia; hyperlipidemia; diabetes; omics; metabolomics; genomics

1. Introduction

Diabetes mellitus (DM) is a multifactorial disease with high mortality worldwide. Chronic DM is the eighth-leading cause of deaths globally, responsible for 1.5 million deaths each year [1]. According to the World Health Organization (WHO), in 2013, 381 million adults were diagnosed with DM, which increased to 422 million in 2016 and is expected to double by 2030. Type 1 (T1DM)
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represents 15% of cases, and the remaining cases are type 2 (T2DM) [1]. T2DM is primarily treated with pharmacotherapeutic drugs, evidence-based alternative approaches, and functional food-based approaches [2]. Pharmacotherapeutic approaches generally consist of monotherapy or binary/polytherapy, depending on severity. Most physicians use the binary approach and prescribe insulin-secretogenic sulfonylurea drugs and the insulin sensitivity enhancer metformin. Additional drugs address diabetes-induced vascular complications, with the average number of prescribed daily drugs being as high as four [3]. Combined drug therapy is associated with long-term side-effects and other costs, resulting in non-adherence [4]. Moreover, evidence-based alternative approaches may have safety and toxicity issues due to which precision nutrition-based approaches have recently been proposed as alternatives to defer or deter T2DM and its complications.

The provision of individualized dietary and nutritional recommendations is referred to as precision nutrition. Polyphenol-rich fruits (including berries) are the primary components of precision nutrition, and consumption of these fruits, like berries, represent a potential “frontline strategy” for combating T2DM in obese or overweight patients. Substantial evidence suggests that T2DM onset can be prevented or managed by being regularly consumed berries reduced their T2DM risks by up to 35% [6]. Due to the significance of berry consumption and the lack of comprehensive studies examining berry consumption effects specifically on T2DM, this study aimed to collect and summarize all studies examining the relationship between berry consumption and DM.

DM is a metabolic syndrome with concordance changes in insulin sensitivity and/or availability. This insulin insensitivity and/or deficiency induces derangements in metabolic pathways related to glucose, lipids, and protein metabolism. Berry, or its byproduct, intake not only opposes these derangements by normalizing the metabolic homeostasis of glucose, lipids, and protein metabolism, but also improves insulin sensitivity and secretary indexes. Therefore, all available in vitro and in vivo studies involving whole berries or berry bioproduct consumption and citing the normalization of insulin signaling, secretion, and sensitivity, restoring the altered glucose, lipid, and protein metabolism, and reduction of oxidative stress and inflammatory cytokines were included. In order to determine the hypoglycemic and hypolipidemic potential of berries, studies that added berries to high-fat (HFD) and high-carbohydrate (HCD) diets, defined as diets with >45% fat and >60% carbohydrates, respectively, were also included. In addition to HFD and/or HCD, disruption of intestinal endothelium and homeostasis resulting in epithelial inflammation, increased permeability (i.e., dysbiosis), and alteration in gut microbial taxonomic composition and diversity (increase in Firmicutes:Bacteroidetes ratio, and reduction in intestinal mucosal barrier dysfunction (IMBD) restoring bacterial families, proteolytic and glycolytic microflora, short-chain fatty acids (SCFA), and organic acids (SCOA) producing microflora) are also considered risk factors to obesity and DM. IMBD associated bacterial families protect the epithelial layer of the intestine whereas SCFA and SCOA played important role in the synthesis and production of immunoglobulins and immune-supportive cytokines to protect against dysbiosis and metabolic disorders. In this context, the impact of berry or berry product intake on the attenuation of obesity-associated disorders and dysbiosis was also reviewed. Studies involving the metabolic fingerprinting of berries were also described to represent the possible number of compounds considered responsible for their antioxidative and antidiabetic actions.

Consequently, this review aimed to discuss scientific evidence regarding a positive role of berry consumption on the prevention or delay of diabetes development and reduction or avoidance of diabetes-related complications. Moreover, a detailed composition of different berries is also presented.
2. Methods

Studies examining berry consumption and T2DM were searched for (last time accessed 15 June 2020) in the Medline/PubMed, ScienceDirect/Scopus, and Web of Sciences databases using the following keywords and phrases: berry consumption and diabetes, berry polyphenolic compounds and diabetes, berry intake and glucose metabolism, berries and high-fat diets, berries and high-glycemic diets, metabolic fingerprinting of berries, lipid metabolism and berries, glycomet control, human clinical trials with berries, in vitro/in vivo studies using berries, and individual berry names. The search using these keywords and phrases resulted in more than 3000 articles in said scientific databases, as illustrated in detail in Figure 1. All articles not in line with the objectives of this review article were not considered. Additionally, the articles that were found more than once in these databases were counted once, and after removal of these duplicate and irrelevant records, nearly 2645 publications were thoroughly screened for inclusion eligibility. Finally, 336 publications were found relevant and fit to be reviewed. Only studies examining berries or berry product consumption relative to metabolic syndrome conditions or otherwise DM respective and berry fingerprinting were included. The schematic flow diagram for the selection of studies in this work is presented in Figure 1.

3. Blueberries

Blueberries (BlBs) top the list of five fruits recommended by the Food and Agriculture Organization of the United Nations (FAO) against diabetes, cancer, liver disease, anemia, and cardiovascular disease (CVD). Initially, the in vitro antidiabetic activity of BlBs were reported by Barberis et al. [9] and Martineau et al. [10]. Barberis et al. described the reduced amount of glucose absorption in the Caco-2TC7 monolayer human intestinal cell line in the supplementation of phosphate-buffered-saline (PBS) containing BlB juice (BlBJ) prior to glucose stimulation. Martineau et al. [10] used insulin-dependent/independent 3T3-L1, C2C12, and TC-tet cell lines. The overnight incubation of these cells with BlB extracts (BlBEs) enhanced glucose uptake even in the absence of insulin compared to the vehicle-delivered control cell cultures [10]. The basal secretion of insulin from TC-tet cells increased 2.5 times to 7.5 times with increasing glucose amounts from 6 mM to 10 mM. A significant increase in glucose-stimulated insulin secretion (GSIS) was also seen after treating cells with BlBEs [10]. The BIBE adipogenic effects were also examined by assessing lipid formation and
accumulation in pre-adipocytes, and BIB treatment was almost as effective as the positive control, rosiglitazone, for lipid accumulation. BIB consumption downregulated the HFD-induced upregulation of total cholesterol (TC), low-density lipoprotein cholesterol (LDL-C), leptin, and inflammatory genes (L-6, TNF-α, inducible nitric oxide synthase (iNOS)), monocyte chemo-attractant protein-1 (MCP-1) (an inflammatory cytokine), peroxisome proliferator-activated receptors γ (PPAR-γ), and fatty acid synthase (FAS) [11,12].

Hypoglycemic and hypolipidemic potential of BIBs or its polyphenol rich products has also been checked in many in vivo studies; Grace et al. [13] fed streptozotocin (STZ)-induced diabetic rats diets supplemented with phenolic- and anthocyanin-enriched BIBES. Anthocyanin-enriched diets increased hypoglycemic activity (51%) compared with phenolic-enriched diets (33%) and metformin-treated controls (32%), suggesting that anthocyanins modulated hyperglycemic and hyperlipidemic activities [13]. The supplementation of BIBE increased the beneficial glucose metabolism involved peroxisome proliferator response element (PPRE) (1.3–1.8%), glucose transporter 2 (GLUT-2) (1.5%), and PPAR-γ (1.4%) activities, and reduced the proinflammatory nuclear factor (NF)-κB activity [14]. Furthermore, an increase in the intercellular levels of the mRNA of glucose transporter (GLUT4), insulin receptor substrate-1/2 (IRS-1/IRS-2) (insulin response mediators regarding glucose metabolism), and AMP-activated protein kinase (AMPK) (a key regulator of mitochondrial biogenesis and cellular energy homeostasis) were observed in skeletal muscles, indicating increased glucose uptake [15,16]. BIB metabolites, especially anthocyanins, also promoted glucagon-like peptide-1 (GLP-1) expression and PPAR activity; GLP-1 increases glucose-dependent insulin secretion and pancreatic β-cell proliferation, whereas PPAR and nuclear fatty acid receptors improve IR [17]. Few studies have also shown improved insulin resistance (IR) but with conflicting results in reduced BW gains [16,18,19]. However, in human clinical trials, improved insulin sensitivity without BW changes was observed [20]. Thus, insulin sensitivity may increase following BIB consumption, but BIBE may be less effective for modulating weight loss. Besides the BIBE, a few studies have also used the whole BIB fruit to determine its antidiabetic and anti-obesity potential in a group of people at high risk of T2DM (Table 1). BIB smoothie supplementation significantly reduced insulin resistance (IR) [21]. The ad libitum consumption of 100% pure BIB showed hypoglycemic activity, suppressing fatty acid synthase (FAS)- and β-oxidation-related gene expression in HFD-fed C57BL/6 mice (Table 1) [12]. Alcohol-free fermented juice, containing 30% BIBJ and 70% blackberry juice, reduced epididymal fat pad weights, percent fat mass, plasma triglyceride, and total cholesterol (TC) levels as well as mean adipocyte diameters and improved fasting blood glucose and GTT levels [22]. In another study, BIBJ consumption increased glucose uptake and inhibited adipogenesis by reducing adiponectin levels in KKKy mice [23]. In addition to BIBJ, BIB powder consumption in sugar-matched/sugar-non-matched smoothies extended the post-prandial glucose response and reduced peak postprandial glucose levels [24]. Diabetes and obesity are inter-linked via chronic inflammatory conditions, where macrophages infiltrate and accumulate in adipose tissue, triggering pro-inflammatory cytokine secretion [25]. BIB supplementation reduced these (pro)-inflammatory cytokine secretions (i.e., NF-κB, interleukin (IL)-10, tumor necrosis factor (TNF)-α, and IL-6 expression) in obese and diabetic mice [25]. BIBE consumption also showed excellent anti-inflammatory effects against soluble vascular cell adhesion molecule-1 (sVCAM-1) (inflammatory biomarker), MCP-1, C-reactive protein (CRP) (acute inflammatory protein), and vascular endothelium [26].

Oxidative stress increases reactive oxygen species (ROS), chemokines, nitric oxides (NOs), adhesion molecules, nuclear factor (IkBα) production, and glycation prior or after diabetes. Human aortic endothelial cells (HAECs) treated with purified BIB anthocyanins (hippuric acid, hydroxyhippuric acid, isovanillic acid-3-sulfate, benzoic acid-4-sulfate, and vanillic acid-4-sulfate) demonstrated reduced ROS, chemokine, NO, adhesion molecule, and IkBα production [27]. In a human clinical trial, post-exercise blueberry BIB consumption decreased manganese superoxide dismutase (Mn-SOD) levels [28]. Li et al. [29] reported anti-oxidative and anti-inflammatory cytokine marker suppression by 19 and 31%, respectively, in adipocytes and macrophages co-cultured with piceatannol, a BIB-derived bioactive compound. Piceatannol also ameliorated malfunctioning
insulin-stimulated glucose uptake by upregulating Akt phosphorylation (crucial for IRS activation and hence increasing insulin sensitivity) and forkhead box O1 (FOXO1) (a transcription factor). Pterostilbene, a PPAR-α agonist found in BlB, promoted fatty acid catabolism by upregulating (up to 3%) of AMPK, carnitine palmitoyl transferase-1 (CPT-1) (an enzyme for long-chain fatty acid-LCFA oxidation), acyl-CoA oxidase (ACOX) (enzyme of β-oxidation system), and uncoupling protein-2 (UCP-2) (a protein involved in glucose disposal, insulin secretion, and cellular energy dissipation) expression. AMPK is associated with mitochondrial energy production, and AMPK activation regulates liver cell gluconeogenesis by suppressing glucose-6-phosphatase (G6Pase), phosphoenolpyruvate carboxykinase (PEPCK) (gene involved in glyceroneogenesis and gluconeogenesis), FOXO1, PPAR-γ coactivator 1α (PGC-1α), and glucose production. FOXO1 regulates PEPCK, PGC-1α, and G6Pase expression, thus affecting glucose release [30].

The integration of BlB polyphenols with a HFD also attenuated HFD rendered disorders and dysbiosis. The BlB powder supplementation improved the systematic inflammation and insulin sensitivity by modulating the gut microbial population in rat fed on a HFD [31]. In human, BlB intervention offered the prebiotic-effect by increasing the relative abundance of beneficial fermentative bacterium Bifidobacterium spp [32]. The BlB-derived anthocyanins also improved the IMBD restoration by decreasing the population of E. coli [33]. More recently, Rodríguez-Daza and Daoust [34] also witnessed that BlB-derived proanthocyanidins did not significantly improve the dysbiosis symbolic Firmicutes:Bacteroidetes ratio, but its supplementation did improve the population of genera (Akkermansia, Adlercreutzia, an unknown genus of order Clostridiales, Peptostreptococcaceae, and Ruminococcaceae) considered responsible for the maintenance and restoration of the colon mucosal barrier. The health promoting role of BlB and its byproducts can be explained further by a comprehensive metabolite profile for BlB/BIBE [10,35,36,37,38,39] and is shown in Table 2. The metabolic fingerprinting reveals BlB/BIBE as a rich source of antioxidative, antidiabetic, anti-inflammatory anthocyanins, proanthocyanidins, epicatechins, aglycons, glycosides, catechins, phenolic acids, chlorogenic acids, caffeic acid derivatives, and quercetin derivatives. Collectively, these studies demonstrated that BlB supplementation protected against HFD/HCD-induced IR hyperglycemia, pro-inflammatory responses, oxidative stress, adipocyte death, and improved insulin sensitivity, with mixed results for HFD-induced BW gain. The identified anthocyanins associated with these activities include glucosides, galactosides, and arabinosides of cyanidin, delphinidin, malvidin, peonidin, and petunidin.
Table 1 A comprehensive list of berry interventions and their consequent effects on various levels.

| No. | Study Design | Study Subject | Duration | Berry Interventions | Intervention Diet | Significant Findings | Ref. |
|-----|--------------|---------------|----------|---------------------|-------------------|----------------------|------|
| 1   | RCT          | C57bl/6j mice (n = NS) | 12 wk | LFD (20% kcal from lard fat) and HFD (70% kcal from lard fat) | Lower the blood glucose level and dyslipidemia markers | [13] |
| 2   | RCT          | Male C57BL/6 mice (n = 24) | 8 wk | 4% (w/w) F/D whole BIB powder with HFD provided ad libitum | HFD (60% kcal from fat) | Offer protection against HFD-induced obesity, adipose tissue macrophages inflammatory gene expression, and oxidative stress | [11] |
| 3   | RCT          | C57/B6 mice (n = 200) | 12 wk | 5% or 10% (w/w) of whole BIB with HFD provided ad libitum | HFD (45% kcal from fat) | Reduced HFD-induced cellular inflammatory cytokines, chemokines, interleukins, and proinflammatory interferon gamma-producing T-cells | [18] |
| 4   | RCT          | Male Zucker Fatty and Zucker Lean rats (n = 48) | 8 wk | 4% (w/w) F/D whole BIB powder with HFD provided ad libitum | HFD (45% kcal from fat) | Hypolipidemic, Hyper-insulinemic, hypoglycemic and anti-inflammatory | [40] |
| 5   | RCT          | C57BL/6 mice (n = 48) | 12 wk | Ad libitum 100% BIBJ with HFD provided ad libitum | LFD (20% kcal from fat) and HFD (45% kcal from fat) | Reduced expressions of inflammatory and FA synthesis genes. Reduced IR and plasma dyslipidemia markers | [12] |
| 6   | RCT          | C57BL/6 mice (n = 72) | 8 wk | 65.1 ± 1.6 mg cyanidin-3-O-glucoside/L (from 30% BIB + 70% blackberry juice available ad libitum) | LFD (60% kcal from fat) | Anti-obesity, hypoglycemic, antidiabetic | [22] |
| 7   | RCT          | C57BL/6 and KKAY mice (n = 20) | 4 wk | BIBJ (40–80 mL/kg per day in drinking water) | Normal chow diet | Improved glucose tolerance, reduced glycemic response suggesting increased insulin sensitivity | [23] |
| 8   | RCT          | Obese Zucker rat (n = 20) | 8 wk | 8% wild BIB diet (WB) provided ad libitum | NA | Downregulated expression and plasma concentrations of NF-kB, TNFα, IL-6, CRP in liver and adipose tissues | [25] |
| 9   | SB and RCT   | Obese men and women (n = 66) | 8 wk | 50 g/FDBB per day | NA | Reduction in plasma oxidized LDL and other plasma lipid oxidation products | [20] |
| 10  | DB, PC, RCT  | Overweight or obese individuals (n = 30) | 4 wk | 4 g of insulin/day from BIB (equivalent to two cups of whole BIB) | NA | Improvement in glycemic response, insulin sensitivity, satiety, serum lipid parameters, and fecal markers of gut microbiota | [41] |
| 11  | DB, PC, RCT  | Diabetic patients (n = 58) | 24 wk | 160 mg of BIB anthocyanins twice daily | NA | Reduced serum concentration of LDL-C, TG, apolipoprotein, apolipoprotein C-III, lipid and protein oxidation markers with strengthening the inherent antioxidative system | [42] |
| 12  | DB, PC, RCT  | Healthy adults (n = 44) | 6 wk | 45 g/day F/D BIB powder | 12-oz yogurt and skim milk-based smoothie | Improvement in endothelial function in subjects with metabolic syndrome | [21] |
| 13  | DB, CO, RCT, | Healthy human beings (n = 17) | 4 wk | 310–724 mg/kg/BW.day BIBanthocyanin | Sugar-matched smoothie | Extend the postprandial glucose response beyond the period observed for a sugar-matched control | [24] |

(1) Blueberries (BIB) (Animal studies)
| Study | Design | Species | Treatment | Duration | Intervention | Outcome |
|-------|--------|---------|-----------|----------|--------------|---------|
| 1     | RCT    | Male KK-Ay mice (n = 16) | 5 wk | 27 g of BB extract/kg diet daily | NA | Activation of AMP-activated protein kinase (AMPK) resulting in increased insulin sensitivity, upregulation of glucose transporter GLUT4, suppression of glucose production in liver |
| 2     | RCT    | Male Fischer rats (n = 24) | 16 m | 2% whole CrB powder standard NIH-31 rodent chow available ad libitum | NA | Increased β-cell glucose responsiveness; age related decline in in basal plasma insulin concentrations was delayed by cranberry |
| 3     | RCT    | Brown Norway (BN) rats (n = 96) | 6 wk | BB extract 100 mg/kg BW/day | Normal chow diet | Prevent diabetic retinopathy |
| 4     | RCT    | Male KM mice (n = 60) | 5 d | BB extract (containing 42.04% anthocyanins) 200 mg/kg BW-day | Normal chow diet | Reduced β-cell glucose responsiveness; age related decline in in basal plasma insulin concentrations was delayed by cranberry |
| 5     | RCT    | Goto-Kakizaki (GK) rat (n = NS) | 4 wk | BB decoction with rodent chow | Powdered rodent chow | | (Human studies) |
| 6     | RCT    | Healthy men and women (n = 9) | 1 d | 10% BB in fermented drink up to 300 g/day | White bread | Anti-inflammatory markers, vascular health markers and reducing capacity |
| 7     | DB, CO, RCT | T2DM Male volunteer subjects (n = 8) | 24 h | A single capsule of 0.47 g BB extract (36% anthocyanins) | NA | Decrease in the incremental AUC for both glucose and insulin without alterations in GLP1, glucagon, amylin, and anti-inflammatory peptides |
| 8     | CO, DB, RCT design | Obese/Overweight/diabetic men and women (n = 16) | 3 wk | 3 x 0.47 g of Mirtoselect capsules per day, a standardized BB extract (36% anthocyanins) | NA | Reduced activity of digestion enzymes without alterations in anti-inflammatory markers, vascular health markers and reducing capacity |
| 9     | RCT    | Healthy men and women (n = 62) | 4 wk | BB juice 330 mL/day | NA | Anti-inflammatory markers, vascular health markers and reducing capacity |
| 10    | RCT    | Healthy men (n = 40) | 8 wk | Fresh BB 100 g/day of BB | NA | | (I) Cranberries (CrB) | (Animal studies) |
| 11    | Parallel RCT | Healthy men and women (n = 27) | 8 wk | Fresh BB 400 g/day | NA | Reduction in the low-grade inflammation with different cytoplasmic ribosomal proteins, Toll-like receptor, and B-cell receptor signaling pathways |
| 12    | RCT    | Healthy men and women (n = 9) | 1 d | 10% BB in fermented drink up to 300 g/day | White bread | Anti-inflammatory markers, vascular health markers and reducing capacity |

**Notes:**
- **HFD (High-Fat Diet):** 65% lipids, 15% proteins and 20% carbohydrates.
- **ADI (Ad libitum):** Food and water were available ad libitum.
- **HOMAIR:** Homeostasis model assessment of insulin resistance.
- **ALT:** Alanine aminotransferase.
- **MDA:** Malondialdehyde.
- **NO:** Nitric oxide.
- **GLP1:** Glucagon-like peptide 1.
- **TG:** Triglycerides.
| No. | Design | Participants/Group | Duration | Intervention | Outcome Measures |
|-----|--------|---------------------|----------|--------------|------------------|
| 3   | CO, RCT | Obese participants (n = 25) | 2–4 h | Cranberries (40 g) | HF breakfast (70 g fat, 974 kcal) | Improved postprandial glycemic control, reduction in lipid oxidation products and inflammatory cytokines [56] |
| 4   | PC, DB, RCT | T2DM men and women (n = 30) | 12 wk | Dextrose sweetened normal calorie CrB juice (NCCBJ) [27% CB]; 27% CrB; 190 Cal/240 mL | 5 g Vanilla Crisp Power Bar (contained 230 Cal, 2.5 g total fat, 3 g dietary fiber, 20 g sugars, 22 g other carbohydrates, and 9 g protein) | Decrease in the TC: HDL-C ratio [57] |
| 5   | single CO RCT | Healthy men and women (n = 12) | OTCS | 3 capsules of CrB extract/day (1 capsule = 500 mg) | NA | Improved metabolic response towards glucose [58] |
| 6   | RCT | Non-diabetic men and women (n = 187) | OTCS | 200 x 2 mL RCCJ was enriched with omega-3 fat acid (180 mg EPA + 120 mg DHA) on daily basis | NA | Antioxidative defense system [68] |
| 7   | RCT | T2DM men and women (n = 13) | OTCS | Raw CrB (55 g, 21 cal, 1 g fiber); Sweetened dried CrB (40 g, 138 cal, 2.1 g fiber); Sweetened dried CrB containing less sugar (SDC-LS) | White bread (57 g, 160 cal, 1 g fiber) | Favorable glycemic and insulinemic response [60] |
| 8   | CS Nutrition Examination Survey (n = 10 891) | Healthy men and women | 2 days | Average 2-day CrB intake 158 to 404 mL | Routine diet | Lowered the weight-gain, TC, and proinflammatory serum CRP levels [61] |
| 9   | DB, CO, RCT | Healthy men and women (n = 12) | OTCS | 37.5 g of CrB in addition to 37.5 g x 3 of other berries (bilberries, strawberries, blueberries) + 35 g added sugar | NA | Hypoglycemic and hypo-insulinemic [62] |
| 10  | Parallel RCT | Diabetic men and women (n = 48) | 8 wk | Omega-3 fatty acid (180 mg EPA + 120 mg DHA) on daily basis | usual diet and physical activity during the study | Anti-dyslipidemic and hypoglycemic [63] |
| 11  | Parallel DB RCT | T2DM male patients (n = 58) | 12 wk | 1 cup (240 mL) CrB juice daily | NA | Antiglycation, antidabetic, reducing CVD risk factors in T2DM male patients [64] |
| 12  | Parallel DB, PC RCT | Healthy men and women (n = 56) | 8 wk | 480 mL (80 kcal) whole CrB juice daily | American foods (HFD) and 3–5 servings of fruits or vegetables daily (328–618 g/d depending on energy intake) | Anti-dyslipidemic, hypoglycemic, improved HOAM-IR [65] |
| 13  | RCT | Patients with metabolic syndrome (n = 55) | 60 d | 0.7 L/day (120 kcal) of reduced-energy CrB juice containing 66 mg proanthocyanidins; total phenolics of 104 and 0.12 mg folic acid | NA | An increase in adiponectin and folic acid and a decrease in homocysteine, decreased lipoperoxidation and protein oxidation levels [66] |
| 14  | CO, DB | Obese yet healthy men (n = 35) | 4 wk | Increasing doses of low-calories CrB during three successive periods of 4 wk (wk 1–4: 125 mL/day, wk 5–8: 250 mL/day, and wk 9–12: 500 mL/day) | NA | Improved augmentation in obese men [67] |
| 15  | DB, CO | Obese men (n = 30) | 12 wk | NA | Improved antioxidative defense system [68] |
| Study | Design | Participants | Intervention | Outcome Measures |
|-------|--------|--------------|--------------|------------------|
| 16    | DB, CO | Obese men (n = 30) | Increasing doses of low-calories CrBJ during three successive periods of 4 wk (wk 1–4: 125 mL/day, wk 5–8: 250 mL/day, and wk 9–12: 500 mL/day) | Decrease in plasma OxLDL, intercellular adhesion molecule-1 (ICAM-1), vascular cell adhesion molecule-1 (VCAM-1) and E-selectin concentrations [69] |
| 17    | CO, DB | Abdominally obese men (n = 30) | Increasing doses of low-calories CrBJ during three successive periods of 4 wk (wk 1–4: 125 mL/day, wk 5–8: 250 mL/day, and wk 9–12: 500 mL/day) | Increased plasma HDL-cholesterol concentrations [70] |
| 18    | RCT    | Healthy men (n = 21) | CrBJ 7 mL/kg,BW.day (i) Raspberries (RB) (Animal studies) | Reduction in plasma OxLDL levels and Improved antioxidative defense system [71] |
| 1     | RCT    | Weanling male Syrian golden hamsters | RBJ 275 mL/day (1 mL = 0.6 g of berries) semi-purified hyperlipidic diet (0.5% cholesterol and 15% lard) | Hypo-cholesterolemic and antioxidative [72] |
| 2     | RCT    | Male Wistar rats (n = 30) | Dose of ellagitannins enriched RB extracts equivalent to daily consumption of 125 g of fresh fruit by a human healthy adult of 70 kg (i.e., 20 mg/kg BW.day orally) | Protection from the ethanol induced oxidative stress and inflammatory biomarkers [73] |
| 3     | RCT    | Male Lewis rats (n = 24) | RB extracts at 30-120 mg/kg,BW | Inhibition of inflammation, pannus formation, cartilage damage, and bone resorption [74] |
| 4     | RCT    | CD1 male mice (n = 36) | RB infusion by gavage (100 mg/kg BW.day) 5.3% RB supplementation along agar-based diet finally containing polyphenolics (963 mg extractable GAE/kg agar-based diet) | Improved antioxidative defense system [75] |
| 5     | RCT    | Obese diabetic (db/db) mice (n = 30) | 5 mL/kg BW.day | Hyper-cholesterolemic and diabetes-induced oxidative stress [76] |
| 6     | RCT    | Male Zucker Fatty rats (n = NS) | 20 g of diet per day containing RB (2% red raspberry F/D powder) | Upregulation of the expression of myocardial adiponectin receptor 1 and apolipoprotein E, improving the plasma cholesterol and triglyceride homeostasis [77] |
| 7     | RCT    | Male Wistar rats (n = 42) | 313 g whole RB with/without *Lactobacillus plantarum* HEAL19 (HEAL19 at 10⁵ cfu) per day with diet | Increased intestinal SCFA load and anti-inflammatory [78] |
| 8     | RCT    | Male F-344 rats (n = NS) | AIN-76A diet containing either 5% whole BRB powder, 0.2% BRB anthocyanins, or 2.25% of the residue fraction provided ad libitum | Anti-dysbiosis, anti-inflammatory, anti-obesity [79] |
| 9     | RCT    | Male db/db mice with C57BL/6j Background (n = 48) | 150 mg/kg BW.day per mice RB derived pelargonidin-3-O-glucoside | Hypoglycemic, anti-inflammatory, anti-obesity [80] |
| 10    | RCT    | Specific-pathogen free C57BL/6 mice (n = 20) | AIN-76A diet with 10% black raspberry powder provided ad libitum | Hypo-glycemic, anti-metabolic syndromic [81] |
| 11    | RCT    | Male db/db mice (n = 30) | 10% F/D RB in a isocaloric standard diet | Hypo-cholesterolemic, antioxidative, improved insulin sensitivity [82] |
| 12    | RCT    | C57BL/6j mice (n = NS) | Energy-containing RB foods (puree and puree concentrate and whole fruit powder) containing 10% raspberry and HFD supplemented with 0.2% (w/w) RB extract provided ad libitum | Anti-obesity and antidiabetic [83] |
| Study 1 | Design | Populations | Duration | Intervention | Comparison | Outcomes |
|---------|--------|-------------|----------|--------------|------------|----------|
| 13      | RCT    | C57BL/6j, C57BL/Ks db/db, and db/+ male mice (n = NS) | 8 wk | 0.2% Cyanidin 3-glucoside in HFD | HFD (58% of calories from coconut hydrogenated fat) | Anti-obesity, anti-inflammatory, improvement in the insulin sensitivity |
| 14      | RCT    | Male Sprague Dawley rats (n = 40) | 8 wk | Application of RB derived EA (0.1–10 mg/mL) on ischemic stomach (1.5 mL/100 g,BW) in an in an ex vivo chamber | NA | Gastric protective action against gastric lesions induced by NH4OH, due to anti-oxidative activity of EA |
| 15      | RCT    | Male Wistar rats (n = 22) | 4 wk | Oral administration of 10–20 mg/kg,BW of RB derived ellagic acid | NA | Anti-inflammatory and anti-oxidative reduced ectopic lipid storage, alleviated inflammation responses, improved whole-body insulin sensitivity, and promoted mitochondrial biogenesis |
| 16      | RCT    | Male Wistar rats AMPKα1−/− (n = 12) | 10 wk | 5% supplementation of RB extracts (contains polyphenols at ~11 g gallic acid equivalent (GAE)/(g of DW) along HFD | HFD (60% from fat) | Anti-dyslipidemic, hypoglycemic |
| 17      | RCT    | Male mice (C57BL/6) (n = 40) | 12 wk | 5% F/D RB powder in HFD provided ad libitum | HFD (60% energy from fat) | Anti-obesity, anti-inflammatory improvement in the insulin sensitivity Anti-dyslipidemic, hypoglycemic, attenuated hepatic ER and oxidative stresses, as well as adipocyte inflammation |
| 18      | RCT    | Male KK-Ay mice (n = NS) | 5 weeks | Cynanidin 3-glucoside 2 g/kg,BW.day in the normal chow diet | HFD + High-sucrose diet (37% energy from sucrose) | Reduced the postprandial insulin response, improved the glycemic profile, improved postprandial glucose metabolism. |
| 19      | RCT    | Male mice (C57BL/6) (n = 40) | 12 wk | 3% RB seed flour (equivalent to 0.03% ellagic acid) in HFD and HFD + High-sucrose diet | HFD (41% energy from fat) | Anti-dyslipidemic, anti-inflammatory, anti- obesity |
| 20      | PC, CO, RCT | Healthy men and women (n = 20) | 4 wk | Human studies | Macronutrient-matched high-carbohydrate cereal bars (45% total sugars) | Anti-dyslipidemic, hypoglycemic, attenuated hepatic ER and oxidative stresses, as well as adipocyte inflammation |
| 21      | RCT    | Healthy men and women (n = 12) | NS | 100 g RB along the designated diet | High-carbohydrate food in the form of pancakes (50 g available carbohydrate from 333 kcal pancake) | Postprandial hyperglycemia to sustainable glycemic response |
| 22      | 3 randomized, controlled, CO, | Healthy women (n = 13-20) | OTCS | 150 g whole berries puree along each meal study 1: white bread + strawberries, bilberries, or lingonberries study 2: white bread + h raspberries, cloudberris, or chokeberries study 3: white bread or rye bread + mix berries consisting of equal amounts of strawberries, bilberries, cranberries, and blackcurrants | White bread or rye bread with 50 g available starch | Reduced the postprandial insulin response, improved the glycemic profile, improved postprandial glucose metabolism. |
| 23      | CO, RCT | T2DM men and women (n = NS) | 12 wk | 250 g frozen red raspberries puree with each breakfast (l) Mulberries (MBs) (Animal studies) | NA | Anti-dyslipidemic, anti-inflammatory, anti-obesity |

### Control Groups

| Study 1 | Design | Populations | Duration | Intervention | Comparison | Outcomes |
|---------|--------|-------------|----------|--------------|------------|----------|
| 1       | Randomized block design | Male C57BL/6 mice (n = 60) | 8 wk | MB anthocyanins at 200 mg/kg HFD provided ad libitum | HFD (45% kcal from fat) | Anti-dyslipidemia, anti-inflammatory, anti-obesity |
| 2       | RCT    | Male db/db mice with C57BL6/J genetic background (n = 50) | 8 wk | MB fruit extracts 50 and 125 mg/kg BW every day orally by gavage | NA | Antioxidative and hypoglycemic |
| 3       | RCT    | male adult Wistar rats (n = 70) | 6 wk | MB fruit wine 400 mL/70 kg of body weight daily | NA | Antioxidative and hypoglycemic |
| 4       | RCT    | male Sprague-Dawley rats (n = 50) | 8 wk | MB fruit derived cyanidin-3-O-β-D-glucopyranoside (10 mg/kg,BW, daily) orally by gavage | NA | Antidiabetic cystopathy |
| Study | Design | Subject Details | Treatment | Outcome Details |
|-------|--------|-----------------|-----------|----------------|
| 5     | RCT    | Adult diabetic male Wistar rats (n = 12) | 6 wk MB polysaccharides (200 mg/kg BW daily) in HFD provided ad libitum | HFD (1% cholesterol and 10% corn oil) improved oral glucose tolerance/insulin resistance, bioactivities of superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GPx), were increased |
| 6     | RCT    | Male Gold Syrian hamsters (n = NS) | 12 wk Water extracts of MB fruit at 1–2% (w/w) in HFD provided ad libitum | HFD (45 kcal/kg) Hypolipidemic effects |
| 7     | RCT    | Male C57BL/6 mice (n = 48) | 12 wk Anthocyanin from MB of 40-200 mg/kg of HFD | HFD (1% cholesterol and 10% corn oil) improved oral glucose tolerance/insulin resistance, bioactivities of superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GPx), were increased |
| 8     | RCT    | Male Syrian golden hamsters (n = 32) | 12 wk Water extracts of MB fruit at 0.5–2% (w/w) in HFD provided ad libitum | HFD (1% cholesterol and 10% corn oil + 0.1% cholesterol) improved oral glucose tolerance/insulin resistance, bioactivities of superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GPx), were increased |
| 9     | RCT    | Male Wistar rats (n = 32) | 4 wk 5–10% (w/w) mulberry fruit polysaccharide fractions in HFD provided ad libitum | HFD (1% cholesterol, 0.5% sodium cholate, and 20% commercial diet) improved oral glucose tolerance/insulin resistance, bioactivities of superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GPx), were increased |
| 10    | RCT    | Male C57BL/6 mice (n = 48) | 6 wk 0.5–2% (w/w) water extracts of MB fruit in high-fat (35% kcal from fat) ethanol rich liquid diet (36% kcal from ethanol) provided ad libitum | HFD (1% cholesterol, 18% lipid (lard), 40% sucrose) Anti-obesity, hypoglycemic, antioxidative, anti-inflammatory effects |
| 11    | RCT    | Male Sprague-Dawley rats (n = 40) | 10 wk MB fruit extracts 100 or 200 mg/kg BW day | HFD (1% cholesterol, 18% lipid (lard), 40% sucrose) Hypolipidemic and improved the enzymatic antioxidant system |
| 12    | RCT    | Female Wistar rats (n = 48) | 20 wk Microencapsulated 50 to 250 mg/kg BW day mulberry fruit extracts (microencapsulated) with HCHF diet which contained total energy around 4.62 kcal/g (fat 31.54%, protein 20.25%, and carbohydrate 48.21%). | HFD (1% cholesterol, 18% lipid (lard), 40% sucrose) Anti-obesity, hypoglycemic, antioxidative, anti-inflammatory effects |
| 13    | RCT    | Male, C57BL/6f mice (n = 12) | 13 wk 20% MB powder in HFD provided ad libitum | HFD, 60% calories from fat Anti-obesity, antidiabetic, increase of Bacteroidetes/Firmicutes ratio |
| 14    | RCT    | db/db mice (n = 50) | NS MB polysaccharide fractions (200–800 mg/kg BW) | HFD, 60% calories from fat Anti-obesity, antidiabetic, increase of Bacteroidetes/Firmicutes ratio |
| 15    | RCT    | Male C57BL6f genetic background (db/db) mice (n = 60) | 8 wk Mulberry fruit extract 25–250 mg/kg BW daily | HFD (35% fat, 20% protein, and 36.5% carbohydrate) Attenuates hepatic steatosis hyperglycemia, hyperlipidemia. Improves insulin signaling and decrease the leptin secretion. |
| 16    | RCT    | Adult diabetic male Wistar (n = 40) | 7 wk MB fruit polysaccharide fractions MFP50 and MFP90 (400 mg/kg BW) (l) Lingonberries (LB) (Animal studies) | HFD (35% fat, 20% protein, and 36.5% carbohydrate) Attenuates hepatic steatosis hyperglycemia, hyperlipidemia. Improves insulin signaling and decrease the leptin secretion. |
| 1     | RCT    | Male C57BL/6 mice (n = NS) | 8 wk LB extracts (125, 250, and 500 mg/kg) in HFD provided ad libitum | HFD (35% fat, 20% protein, and 36.5% carbohydrate) Attenuates hepatic steatosis hyperglycemia, hyperlipidemia. Improves insulin signaling and decrease the leptin secretion. |
| 2     | RCT    | SHR rats (n = NS) | 8 wk Cold-compressed LB juice provided ad libitum | HFD (35% fat, 20% protein, and 36.5% carbohydrate) Attenuates hepatic steatosis hyperglycemia, hyperlipidemia. Improves insulin signaling and decrease the leptin secretion. |
| Study Number | Design | Participants | Study Duration | Intervention | Outcome Measures |
|--------------|--------|--------------|---------------|--------------|------------------|
| 1            | RCT    | Male Wistar rats (n = 32) | 5 wk          | NA           | NA               |
| 2            | RCT    | C57BL/6 mice (n = 60)      | 12 wk         | HFD (45% kcal from fat) | Anti-inflammatory, anti-hypertensive, anti-hypercholesterolemia, antioxidative |
| 3            | RCT    | Male C57BL/6/JBomTac mice (n = 120) | 13 wk         | 20% (w/w) F/D LB in HFD provided ad libitum | HFD (45 kcal% fat) |
| 4            | RCT    | Male C57BL/6/JBomTac mice (n = NS) | 11 wk         | 20% (w/w) F/D LB in HFD provided ad libitum | HFD (45 kcal% fat) |
| 5            | RCT    | Male Apoe/- mice (n = 35)  | 8 wk          | 44% lingonberry + HFD | HFD (38 kcal% fat) |
| 6            | RCT    | Male C57BL/6/JBomTac mice (n = NS) | 11 wk         | 20% (w/w) F/D LB in HFD provided ad libitum | HFD (45 kcal% fat) |
| 7            | RCT    | C57BL/6/JBomTac (n = NS)   | 13 wk         | 20% (w/w) freeze-dried LB + blackcurrants, bilberries or acai berry in HFD provided ad libitum | HFD (45 kcal% fat) |
| 8            | RCT    | Male ApoE/- mice (n = 50)  | 8 wk          | Two LB polysaccharide fractions 15-60 g/kg BW with HFD daily | HFD (38 kcal% fat) |
| 9            | RCT    | Scandinavian type 2 diabetes patients (n = 30) | 12 wk         | Recommended daily intake of LB/berries/fruits | Okinawan-based Nordic diet of about 1,900 kcal/day | Reduced glycemic response, normalized anthropometric parameters |
| 10           | CO, DB, RCT | Healthy normal-weight nonsmoking men (n = NS) | 6 d         | Glycemic diet + 40 g lingonberry powder | Improved metabolic and anthropometric parameters |
| 11           | CO, DB, RCT | Healthy, over-weight, non-smoking male and female volunteers | Single meal challenge | 100 g lingonberry | Nullified the glycemic effect of the sugars present in the meals without affecting the postprandial lipemic response |
| 12           | RCT    | Normal, healthy subjects (n = 9) | 12 wk         | LB polysaccharides + fibers (2 g/Kg of oat bread) | Oat bread |
| 13           | SB, CO, RCT | Healthy women volunteers (n = 20) | 2-h meal tests | Diet 1: 150 g whole LB puree containing 35 g sucrose per day | In reduced glucose and C-peptide response |
| 14           | RCT    | Healthy non-smoking males (n = 14) | 2-h meal tests | Diet 2: 300 mL LB nectar (equal to 150 g fresh berries) containing 35 g sucrose | Optimized postprandial metabolic responses to sucrose with delayed digestion and absorption of sucrose/glucose |

**Notes:**
- RCT: Randomized Controlled Trial
- CO: Controlled Observational Study
- DB: Double-Blind
- HFD: Hypercaloric Meals
- LB: Lingonberry
- BW: Body Weight
- NA: Not Applicable

**Outcome Measures:**
- Significant reduced body fat, lipid accumulation, and plasma levels of the inflammatory marker PAI-1, as well as mediated positive effects on glucose metabolism homeostasis
- Reduced plasma levels of markers of endotoxemia and inflammation
- Decreased triglyceridermia and reduced atherosclerosis
- Improvement in glycaemia, reduction in inflammation and hepatic steatosis
- Downregulation of inflammatory pathways, NF-κB, STAT3 and mTOR as possible targets for antidiabetic therapy
- Hypoglycemic, hypolipidemic, altered caecal microbiota composition
- Improved metabolic and anthropometric parameters

**Diets:**
- Glycemic diet: 200 g yoghurt (lactose-free and fat-free non-flavored natural yoghurt + 50 g glucose)
- Lipemic diet: 200 g Yoghurt (lactose-free and fat-free non-flavored natural yoghurt + 35 canola oil)
- Recommended daily intake of LB/berries/fruits
- Okinawan-based Nordic diet of about 1,900 kcal/day
- Standard diet: white wheat - Hyperlipidic and hypercaloric meals (38 kcal% fat)

**Interventions:**
- Male C57BL/6/JBomTac mice (n = 120)
- Male Apoe/- mice (n = 35)
- Male C57BL/6/JBomTac mice (n = NS)
- Male ApoE/- mice (n = 50)
- Scandinavian type 2 diabetes patients (n = 30)
- Healthy normal-weight nonsmoking men (n = NS)
- Healthy, over-weight, non-smoking male and female volunteers
- Normal, healthy subjects (n = 9)
- Healthy women volunteers (n = 20)
- Healthy non-smoking males (n = 14)
- Male Wistar rats (n = 32)
- C57BL/6 mice (n = 60)
| No. | Design | Group Description | Duration | Intervention | Outcome(s)                                                                 | Reference |
|-----|---------|--------------------|----------|--------------|---------------------------------------------------------------------------|-----------|
| 3   | RCT     | Male DIO C57BL/6j mice (n = 40) | 12 wk    | 6.3%, (w/w) BBR extracts in HFD provided ad libitum | HFD (45% kcal from fat) Anti-obesity, Anti-inflammatory, anti-hypertensive, Anti-obesity, anti-inflammatory, anti-dyslipidemic | [127]     |
| 4   | RCT     | Male Wistar rats (n = 24) | 17 wk    | 25 mg/kg BW BBR extracts in HFD provided ad libitum | HFD (45% kcal from fat) | [128] |
| 5   | RCT     | Male diabetic Sprague-Dawley rats (n = 40) | 40 d | Microfiltrated 12.5-25% BBR juices | Reduced glycaemia (~10.4%), TG (~4.6%) and TC (21.0%), lipid peroxidation, attenuation of oxidative stress | [129]     |
| 6   | RCT     | Male Wistar strain rats (n = 40) | 4 wk     | Normal standard diet with 0.98% BBR polyphenols and 6% BBR fiber | Normal chow diet Anti-inflammatory and anti-dyslipidemic | [130]     |
| 7   | RCT     | Female obese (BKS(D)-Leprdb/J72) and lean (C57BL/6j) mice (n = 24) | 10 wk    | Aged or fresh BBR supplemented at 10% (w/w) of diet provided ad libitum | HFD (60.3% fat, 21.3% carbohydrate and 18.4% protein) Reduced percent fat mass, mean adipocyte diameters, epididymal fat pad weights, and plasma TG and TC. | [22]      |
| 8   | RCT     | Male C57BL/6j mice (n = 72) | 10 wk    | Aqueous, alcoholic and hydro-alcoholic SB extracts (2 g/kg b.w.day) | NA Anti-dyslipidemic | [132]     |
| 9   | RCT     | Diabetic and obese men and women (n = 152) | 1 wk     | Consumption of daily recommended amount of low glycemic index fruit (0.7–1.4 servings/day) | NA Improved plasma and urine antioxidant system | [133]     |
| 10  | open, single-center RCT | Healthy human subjects (n = 6) | 4 h      | 200 mL of BBR juice equivalent to 400 mg of cyanidin equivalent/50 kg of body weight | NA | [134]     |
| 11  | RCT     | Dyslipidemic patients (n = 72) | 8 wk     | 300 mL of BBR juice (equivalent to 316 mg/100 g polyphenols) of BBR with pulp every day | NA Increased apo A-1 and HDL-C along reduction in apo B and hsCRP | [134]     |
| 1   | RCT     | Male diabetic albino Wistar rats (n = 36) | 4 wk     | Aqueous, alcoholic and hydro-alcoholic SB extract (2 g/kg b.w.day) | NA Reduced expression of genes involving glucose, lipid metabolism with improvement in glucose metabolism and liver function Reduction in the oxidative damage in brain and peripheral tissues Reduction in the HFD led increase of FBS, adhesion molecule-1, leptin, E-selectin, resistin, and plasminogen activator protein-1 Improvement of oxidative stress biomarkers, mitochondrial performance, antioxidant enzyme activities, reduction of DNA damage and ROS concentration | [135]     |
| 2   | RCT     | Male Wistar rats (n = 20) | 12 wk    | HFD supplemented with 0.2% irradiated/non-irradiated SB extracts | HFD (47.5% kcal from fat) | [136] |
| 3   | RCT     | Male C57BL/6j mice (n = 36) | 24 wk    | HFD supplemented with 2.6% freeze-dried SB | HFD containing approximately 20% higher in energy density compared to the low-fat diets | [137]     |
| 4   | RCT     | Male Wistar rats (n = 48) | 8 wk     | Supplementation of the diet with a 6% w/w (equivalent to a 5 g/kg 65 BW dose) of a F/D SB-BBB (5:1) powder (FDSSB) | High-fat-sucrose diet (D12451, Research Diet) Anti-obesogenic and anti-inflammatory effects | [138]     |
| 5   | RCT     | Male Wistar rats (n = 24) | 16 wk    | AIN93-modified diet with lyophilized SB extract at 10 g/kg of diet | AIN93-modified diet | [139]     |
| Study ID | Design | Participants | Duration | Intervention | Outcomes |
|---------|--------|--------------|----------|--------------|----------|
| 6 | RCT | Male Wistar rats (n = 20) | 12 wk | Supplementation of 0.2% SB | HFD (47.5% calories from fat) Linseed oil (15 g/day) enriched feed | Antioxidative, anti-stress |
| 7 | RCT | German Landrace pigs (n = 48) | 4 wk | 205-745 g of SB with normal feed per day | Anti-stress and antioxidative |
| 8 | RCT | db/db mice homozygous for the diabetes spontaneous mutation (Leprdb) with C57BL/6 background (n = 24) | 10 wk | 2.35% F/D SB powder in the diet pellets (2/3) (equivalent to two human servings of SB i.e., ~160 g SB) | NA | Increased Bacteriodetes to Firmicutes ratio |
| 9 | RCT | Male CD-1 mice (n = 60) | 8 wk | 5% (w/w) of diet freeze-dried whole SB powder (Human studies) | NA | Increased Bacteriodetes to Firmicutes ratio |
| 10 | DB, RCT, parallel study | Insulin resistant and obese males and females (n = 41) | 6 wk | Beverage containing 1.8 g of a mixture of dry SB and C rP providing 333 mg of polyphenols on daily basis (also equivalent to 112 g consumption of fresh berry fruit) | NA | Improved insulin sensitivity and release |
| 11 | CO, SB, PC, RCT | Hyperlipidemic men and women (n = 24) | 12 wk | SB beverage containing 10 g/serving of freeze-dry SB powder providing 338 mg of polyphenols daily (also equivalent to 110 g consumption of fresh berry fruit) | HFD consisting of typical breakfast food items (i.e., bagel, cream cheese, whole milk, egg, margarine, cantaloupe) | Reduced postprandial lipemia and oxidative stress markers |
| 12 | CO, RCT | Healthy males and females (n = 30) | 5 d | 20 g of five types SB jams each with sugar of different glycemic index | 60 g white bread slice | Non-significant reduction in the postprandial glucose level |
| 13 | CO DB RCT | Healthy males and females (n = 16) | 3 wk | 60 g of three types SB jams each with sugar of different glycemic index and polyphenolic contents | NA | Strawberry jam with high sugar level produced less levels of FFA. |
| 14 | DB RCT | T2DM males and female subjects (n = 36) | 6 wk | Two cups of F/D SB beverage containing 25 g x 2 = 50 g | NA | Reduction in LDL-C and LDL-C/TC and LDL-C/HDL-C ratio |
| 15 | SB, CO parallel, RCT | Obese and overweight men and women (n = 24) | 6 wk | SB beverage containing 10 g/serving of freeze-dry SB powder providing 96 mg of polyphenols on daily (also equivalent to 100 g consumption of fresh berry fruit) | High-carbohydrate-fat diet | Attenuation of diet-induced inflammatory markers |
| 16 | Single-center, CO, SB, PC, | Men and women (n = 26) | OTCS | SB Milk based beverage containing 10 g/305 mL of F/D SB powder | high-carbohydrate, moderate-fat meal (HCFM) | Reduced postprandial insulin and inflammatory response |
| 17 | Four-arm, SB, PC, CO, RCT | Males and females with insulin resistance (n = 23) | NS | SB milkshake containing 10-40 g freeze-dried SB powder where 10 g freeze dried powder = 110 g fresh strawberries | Standard western type meal | Reduced lipid oxidation and post-meal insulin demand |
| 18 | Observatory study | Healthy men and women (n = 247) | 20 years | Dietary flavonoids intake (47-560 mg/day) from fruits and berries | Flavonoid Compounds in Driving Patterns of Microbial Community Assembly | Increased the glutathione level, serum catalase activity, and plasma antioxidant capacity |
| 19 | RCT | Obese men and women (n = 66) | 12 wk | SB beverage containing 25-50 g freeze-dry SB powder daily | HFD (50% calories from fat) | Reduction in the markers of lipid peroxidation (MDA), inflammatory markers (CRP), and reducing trend in HbA1c. |
| 20 | DB RCT | T2DM patients (n = 40) | 6 wk | 50 g of freeze-dried SB powder (equivalent to 500 g fresh strawberries) each day | (I) Goji berries (GB) (Animal studies) | |
|   | Design       | Intervention                                                                 | Duration | Control/Placebo          | Outcome                                                                 |
|---|--------------|------------------------------------------------------------------------------|----------|--------------------------|-------------------------------------------------------------------------|
| 1 | RCT          | Alloxan-induced hyperglycemic/hyperlipidemic adult rabbits (n = 35) and male mice (n = 24) | 10 d     | NA                       | Hypoglycemic and hypolipidemic effect with increased plasma antioxidant capacity [154] |
| 2 | RCT          | Male Wistar rats (n = 70)                                                    | 8 wk     | HFD                      | Significantly reduced liver damage and oxidative changes [155]          |
| 3 | RCT          | Diabetic male mouse of original Kun-ming strain (n = NS)                    | 4 wk     | NA                       | Hypoglycemic and hypolipidemic [156]                                    |
| 4 | RCT          | Obese male Sprague-Dawley rats (n = 60)                                     | 8 wk     | HFD                      | Reduced body-weight-gain with anti-inflammatory properties [157]        |
| 5 | RCT          | STZ-diabetic Male Wistar rats (n = NS)                                       | 8 wk     | NA                       | Increased antioxidative scavenging and antioxidant enzymes. Increased activity of protein kinase C (PKC) [158] |
| 6 | RCT          | STZ-induced diabetic Sprague-Dawley male rats (n = 60)                      | 8 wk     | NA                       | Protective effects in diabetic retinopathy [159]                        |
| 7 | RCT          | Male Wistar rats (n = 16)                                                    | 4 wk     | High-fat-sucrose diet     | Hypoglycemic and improving hyperinsulinemia [160]                        |
| 8 | RCT          | Diabetic male C57BL/6j mice (n = 48)                                         | 7 wk     | HFD                      | Hypoglycemic effects with increased insulin-sensitizing, glucose metabolism, insulin secretion, and promoting pancreatic cell proliferation. [26] |
| 9 | RCT          | Swiss Albino rat (n = 30)                                                   | 3 wk     | NA                       | Hypolipidemic, reduced lipid oxidation, increased insulin-sensitizing and serum antioxidant level [161] |
| 10| RCT          | Diabetic Wistar rats (n = 48)                                                | 8 wk     | HFD and HCD (12% protein, 5% fat, 67% carbohydrate, 5% cholesterol, and 5% other additives) | Reduced serum level of IL-2, IL-6, TNF-α, IFN-α, MCP-1, and ICAM-1 with increased activities of SOD and GSH-Px activities [162] |
| 11| RCT          | Postnatal Royal College of Surgeons (RCS) rats (n = 60)                     | 4 wk     | NA                       | Reduced Caspase-2 activity in experimental group at 25+ post-neonatal day [163] |
| 12| RCT          | Male IL-10-deficient mice (n = 14)                                           | 10 wk    | Normal diet               | Increased gut population of SCFA producing bacteria [164]               |
| 13| RCT          | Kunming mice of clean grade (n = 14)                                         | 2 wk     | Normal diet               | Increased gut population of SCFA producing bacteria, *Firmicutes, Akkermansia, Lactobacillus, and Prevotellaceae* [165] |
| 14| RCT          | Healthy males and females (n = 50)                                           | 30 d     | Intake of 120 mL of GB juice (equivalent to 1632 mg/daily serving 120 mL) of goji berry polyphenols | Increased serum levels of glutathione peroxidase (GSH-Px) and superoxide dismutase (SOD) with reduced level of MDA [166] |
| 15| DB, PC, RCT  | Metabolic syndrome patients (n = NS)                                         | 45 d     | Normal diet               | Reduction in transaminases, waist circumference with improvements in lipid profile, glutathione and catalase level. Increased hepatic antioxidant enzymes, inhibited cytochrome F450 2E1, nitric oxide metabolism and lipid peroxidation [167] |
| 16| RCT          | Male and female C57BL/b6N mice (n = 56)                                      | 8 wk     | GB polysaccharides (1–10 mg/kg BW day) orally | NA [168]                                                                 |
| Study ID | Design | Treatment | Notes |
|---------|--------|-----------|-------|
| 1       | RCT    | Male mice of the C57BL/6 strain (n = NS) | 12 wk | AB seed extracts 300 mg/kg.BW.day by intragastric gavage | HFD (60% calorie from fat) | Reduced expressions of lipogenic proteins (SREBP-1c, pACC, ACC, HMG-CoA reductase) with increased expression of pAMPK, pACC/ACC, and cholesterol transporters (ABCG5 and ABCG8) |
| 2       | RCT    | Zebrafish (n = 70) | 5 wk | HC diet supplemented with10% w/w of AB puree powder | Reduced oxidative markers with lipid lowering effects |
| 3       | RCT    | Oxidatively damaged sod1/sod1 mutant strains Drosophila melanogaster (n = 120) | 5-6d | AB supplemented sugar-yeast (SY) medium to a final concentration of 0.25%, 0.5%, 1% or 2% (w/v) of the food | SY medium | Increased transcript level of gluconeogenesis gene phosphoenolpyruvate carboxykinase (Pepck) with reduction in oxidative stress |
| 4       | RCT    | ApoE-deficient (ApoE 2/2) male mice (n = 23) | 12 wk | F/D açai’ pulp + exercise in progressive treadmill for 30 min daily at a speed of 12 m/min, 0% incline | AIN-93M diet | Hepatic superoxide dismutase activity, mRNA expression of monocyte chemotactic protein-1, percentages of hepatic lipid droplets |
| 5       | RCT    | STZ-induced diabetic Male Wistar rats (n = NS) | 45 d | AB seed extracts 200 mg/kg.BW.day in drinking water | NA | Reduced oxidative damage by reducing the expression of caspase-3, IL-6, TNF-α and MCP-1 |
| 6       | RCT    | Female Fischer rats (n = 32) | 6 wk | Hypercholesterolemic diet (25% soy oil and 1% cholesterol) supplemented with 2% AB (dry wt/wt) | Hypercholesterolemic diet (25% soy oil and 1% cholesterol) | Reduced expression of cholesterol biosynthesis genes HMG CoA-R, EBP-2, ApoB100, LDL-R, ABCG8, and CYPT1A1 |
| 7       | RCT    | STZ-induced diabetic Male Wistar rats (n = 40) | 9 wk | AB seed extracts 200 mg/kg.BW.day by intragastric gavage | HFD (55% calorie from fat) | Hypoglycemic and hypolipidemic with reduced expression of TNF-α and activating the insulin-signaling pathway in muscle and adipose tissue |
| 8       | RCT    | Diabetic female Fisher rats (n = NS) | 30 d | Standard AIN-93 diet supplemented with 2% (w/w) AB pulp | AIN-93 | Modulate ROS production by neutrophils and improve the liver oxidant/antioxidant balance |
| 9       | CO, DB, RCT | Overweight healthy males (n = 23) | Single day meal challenge | Frozen AB pulp (150 g) was prepared in a smoothie with 50 g banana | Lower incremental area under the curve (AUC) for total peroxide oxidative status after açai and increased the iAUC for insulin |

**DB, CO, RCT**

- Healthy overweight men (n = NS)
- Acai berries (AB) (Animal studies)
- No-single-dose-effect on substrate oxidation and prospondial-energy-expenditure

**RCT**

- Male mice of the C57BL/6 strain (n = NS)
- Reduced expressions of lipogenic proteins (SREBP-1c, pACC, ACC, HMG-CoA reductase) with increased expression of pAMPK, pACC/ACC, and cholesterol transporters (ABCG5 and ABCG8)

**CO**

- Healthy overweight males (n = 23)
- Lower incremental area under the curve (AUC) for total peroxide oxidative status after açai and increased the iAUC for insulin
| Study | Design | Participants | Intervention | Outcomes |
|-------|--------|--------------|--------------|----------|
| 10    | RCT    | Male Swiss mice (n = 32) | A single daily dose freeze-dried AB pulp (3 g/kg) via gavage | HFD (32% lard and 1% cholesterol) |
| 11    | four-way CO | Healthy men and women (n = 11) | 100% clarified AB juice/pulp 7 mL/kg BW of each study | NA |
| 12    | open label pilot study | Overweight adults (n = 10) | Intake of 100 g AB pulp twice daily | NA |
| 1     | RCT    | C57BL/6JmsSlc and KK-Ay male mice (N = 10, EACH GROUP) | CB provided ad libitum | Normal chow diet |
| 2     | RCT    | STZ-induced-diabetic-male ICR mice (n = 32) | CB extract (10-100 mg/kg,BW) daily administered orally | NA |
| 3     | RCT    | C57BL/6N mice (n = 20) | CB powder dissolved in water (50 mg/kg daily) | HFD (60 kcal% Lard) |
| 4     | RCT    | Male C57BL/6j (n = 60) | CB extracts (100 mg/kg,BW) dissolved in 0.5% carboxymethyl cellulose | HFD (containing 60% kcal fat) |
| 5     | RCT    | Male Wistar rats (n = NS) | Aronia melanocarpa fruit juice (AMFJ) at doses 10 and 20 mL/kg | NA |
| 6     | RCT    | Polish Merino lambs (n = 24) | 150-300 g of chokeberry pomace per each kg of the complete feed mixture | Complete feed mixture |
| 7     | RCT    | Middle-aged non-medicated subjects with MS (n = 38) an healthy volunteers (n = 14) | CB extracts 100 mg/kg,BW three times daily | NA |
| 8     | RCT    | Male Wistar rats (n = 24) | Diet was supplemented by the extract from CB fruits (0.2% W/W) added at the expense of corn starch | Standard casein diet enriched with 0.5% of cholesterol. Exp group: the diets were modified by 8% of lard and 65% of fructose added at the expense of soybean oil and maize starch, Maltase and sucrase, e improvement of antioxidant status, cholesterol-lowering, |
| 9     | RCT    | Male Wistar rats (n = NS) | CB juice 10 mL/kg,BW.day | NA |
| 10    | RCT    | Male Wistar rats (n = 72) | CB juice 50 mL/kg,BW.day | High-carbohydrate, high-fat + purple maize flour (HPM) |
| 11    | RCT    | Male Wistar rats (n = 36) | CBE at 100 or 200 mg/kg BW.day | Fructose rich diet containing (g/kg diet): casein, 207; DL-methionine, 3-4; fructose, 600; lard, 50; cellulose, 79-8; |
| 12    | RCT    | Male Wistar albino rats (n = 60) | Standardized Aronia extract (SAE) 0.45 mL/kg,BW day) for 4 weeks | HFD (25% fat, 15% protein, 51% starch, and 5% fiber) |

Notes: Nutrients 2020, 12, 2538 | 17 of 66

[179] Attenuated hepatic steatosis and reduced lipid accumulation
[180] Increased plasma antioxidant capacity without affecting generation of reactive oxygen species, and uric acid concentrations in plasma
[181] Postprandial increase in the AUC of plasma glucose with reduced TG, LDL-C, and LDL-C/HDL-C
[182] Duction of glucose-dependent insulinotropic polypeptide (GIP) level
[183] Hypoglycemic, hypolipidemic, antioxidative
[184] Reduced the body and liver weight, lipid accumulation, PPARγ2, FAS, hepatic TG and leptin. Serum transaminases, indicators for liver antioxidant capacity were significantly increased.
[185] Attenuated weight-gain, increase in serum TG, TC, LDL-C and better glucose tolerance
[186] Hypoglycemic, hypolipidemic
[187] Hypoglycemic, hypolipidemic
[188] Beneficial changes in lipid profile, coagulation parameters, inhibition of platelet aggregation
[189] Hypoglycemic, hypolipidemic, antioxidative
[190] Reduced Inflammatory cell infiltration, visceral adiposity index, total body fat mass, improved glucose tolerance
[191] Elevated plasma adiponectin levels and inhibited plasma TNF-α and IL-6. Increased in the expression level of glucose and lipid metabolizing genes
[192] Reduced serum level of TC, TG, LDL-C, with increased serum levels of SFA and PUFA.
| Study Type | Participants | Duration | Intervention | Outcomes |
|------------|--------------|----------|--------------|----------|
| CO open-label trial | T2DM patients (n = 35) | 12 wk | Oral CB juice supplementation (150 mL/day, three times a day for 50 mL) | NA |
| RCT | Healthy female volunteers (n = 29) | 12 wk | 100 mL of polyphenol-rich organic CB juice per day | NA |
| RCT | Apparently healthy women (n = 25) | 12 wk | Consume 100 mL of polyphenol-rich organic CB juice daily | NA |
| RCT | Healthy volunteers and 25 patients with metabolic syndrome (n = 22) | 8 wk | CB extract (3 x 100 mg/day) | NA |
| RCT | Healthy subjects (n = 33) | 4 wk | Consume 200 mL of polyphenol-rich organic CB juice daily (containing 386 ± 9.7 mg of total phenolics expressed as gallic acid equivalents per 100 g) | NA |
| RCT | Diabetic Wistar white male rats (n = 48) | 16 wk | Dose of polyphenols extracts 0.040 g/kg BW every 2 day | NA |
| RCT | Healthy, non-smoking volunteers (n = 11) | 3 wk | CrB juice between meals (250 mL per day) (560 mg GAE/100 mL) | NA |
| RCT | Men with the diagnosed mild hypercholesterolemia (n = 58) | 6 wk | CB juice between meals (250 mL per day) (560 mg GAE/100 mL) | NA |
| 3-arm, DB, parallel RCT | Healthy male volunteers (n = 66) | 12 wk | CB extract capsules (containing 116 mg total polyphenols). CB whole fruit capsules (containing the equivalent to 10 g of the whole CB fruit, and 12 mg of total polyphenols) (1) | NA |

1. **BCE** 5 mg/kg BW (1 mg D3R/kg BW) | Normal diet with IP administration of glucose solution (2 g/kg) | Improved hyperglycemic and hypoinsulinemic condition |
2. **BC** extracts (2 g/kg diet) (equivalent to delphinidine-3-glucoside (D3R) 2 g/kg diet) | NA | Improved glucose tolerance with increased GLP-1 concentration, and upregulation of AMPKα and prohormone convertase 1/3 (GLP-1 precursor) |
3. Diet supplemented with 1% BC powdered extract (32% anthocyanins) | HFD (60 kcal% fat diet) | Protective effect of BC anthocyanins against obesity and associated insulin resistance |
4. HF/HC diet supplemented with 0.1% of BCE (containing 25% anthocyanins and 40% polyphenols) by weight | AIN-93M high fat/high cholesterol (HF/HC) diet (16% fat, 0.25% cholesterol by weight; 55.7%, 125.5% and 31.8% energy from carbohydrate, protein and fat, respectively; 452 kcal/kg) | Reduced BW and adipocyte size of the epididymal fat, energy expenditure and mitochondrial biogenesis genes |
|   | Study Design | Participants | Duration | Intervention | Outcome Measures |
|---|--------------|--------------|----------|--------------|------------------|
| 5 | RCT          | Male New Zealand white rabbits (n = 20) | 4 wk     | Diet supplemented with 1.5% BC polyphenolic extract | Reduced concentration of putrefactive metabolites, β-glucuronidase activity, ameliorated hyperlipidemia, and antioxidative capacity |
| 6 | RCT          | Sprague-Dawley male rats (n = 40) | 4 wk     | 2 mL of BC extract (containing 30 mg BC/kg BW) or 2 mL of CAM30 extract (containing 13.4 mg CAM30/kg body weight), respectively, three times weekly by oral gavage | Reduced β-glucuronidase activity and undesirable bacteria in the caeca. Increased lactobacilli and bifidobacterial gut species |
| 7 | RCT          | Male Sprague-Dawley (SD) rats (n = 40) | 8 wk     | BC extract 100–300 mg/kg BW/day administered orally | Improvements in hypertension, dyslipidemia, insulin resistance, and obesity |
| 8 | RCT          | Male Sprague-Dawley rats (n = 128) | 6 wk     | Diets with dietary fiber and BC extracts (Curtante 30) (containing total anthocyanin 32% (w/w)) | Increased intestinal population of SCFA and total beneficial bacterial population |
| 9 | RCT          | Healthy volunteers (n = 30) | 2 wk     | BC powder CAM30 (672 mg/day; 168 mg x 4 capsules) | Increased intestinal population of SCFA and total beneficial bacterial population |
| 10 | DB, CO, RCT  | Healthy subjects (n = 26) | Single meal challenge test | Standardized high-carbohydrate meal (100 g of white bread) | Reduced Postprandial insulin, C-peptide and GIP, GLUT and SGLT1-mediated glucose transport |
| 11 | DB, CO, RCT  | Healthy subjects (n = 22) | Single meal challenge test | Low-sugar-BC drink containing 300-600 mg anthocyanins | Reduced postprandial insulinemia, glyceria, and incretin secretion |
| 12 | RCT          | Healthy participants (n = 17) | 6 d      | BC powder 6 g/day with water | Improved postprandial AUC of glucose and insulin |
| 13 | DB, CO, RCT  | Endurance-trained females (n = 16) | 7 d      | BC extract 600 mg/day | Increased fat oxidation |
| 14 | RCT          | Healthy sedentary male and female participants (n = 40) | Single meal challenge test | Standardized meal bar to consume for breakfast at least 1 h prior to starting the trial. | Supported positive affective responses |
| 15 | Open exploratory study | Healthy individuals (n = 24) (n = 32) | A single meal challenge study | Two opaque gelatin capsules containing BC anthocyanin (3.2 mg/kg total anthocyanins) | Dose-dependent increase in plasma anthocyanins and recovery from exercise-induced oxidative stress |
| 1   | Pre-diabetic volunteers (n = 43)  | Single dose study | Bearing either 60, 120, or 180 mg Delphinol on each day with one-week washout period | NA | Dose dependently lowered basal insulinemia and glyceria |
| 2   | RCT          | Male balb/c mice (n = NS) | 7 d      | MqB extracts (25, 50 and 100 mg/kg BW) | Ameliorate the oxidative stress condition |
| 3   | RCT          | Male C57BL/BJ mice | 12 wk    | HFD (60% calories from fat) | Decreased glucose production, down-regulation of glucometabolic enzyme |
### Nutrients 2020, 12, 2538

| #  | Study Design | Participants | Intervention | Outcome Measures |
|----|--------------|--------------|--------------|------------------|
| 4  | DB, CO, RCT  | Fifty overweight volunteers (n=42) | 3 capsules of 150 mg standardized maqui berry extract containing 54 mg of anthocyanin daily (equivalent to 162 mg anthocyanins/day) | NA | Reduced levels of Ox-LDL in the anthocyanin group [221] |
| 5  | RCT         | Male C57BL/6Nhsd mice (n=18) | MqB derived Delphinidine (15 mg/kg,BW) daily | High-fat diet and high-carbohydrate drinking water (45% kcal from fat) | Reduced TG accumulation with no effect on metabolic alterations related glucose metabolism [222] |
| 6  | Prospective observational study | Middle-aged participants (n=21) | Two tablets per day of an MCN (Eonlipid) (containing maqui, 300 mg in each tablet) | NA | Improvement of most atherogenesis and oxidative stress markers [223] |
| 7  | CO, RCT     | Healthy male subjects (N=11) | Intake of 250 mL of the MqB drink containing an number of total polyphenols -1000 µmol equivalents of gallic acid | Meals containing food-grade glucose and rice, containing 50 g of carbohydrates by each meal | Reduced glycemic indexed for high-carbohydrate diets. [224] |
| 8  | RCT         | C57BL/6J littermates’ male mice (n=23) | HFD supplemented with 4-5 mg of MqB polyphenols/ 10-15 kcal per day | HFD (45% calories from fat) | Reduced body-weight-gain, improved glucose tolerance and insulin resistance. Differential expression of genes involved in fatty acid oxidation, de novo lipogenesis, thermogenesis, and multilocular lipid droplet formation [225] |

**Note:** Acai berry, AB; AB juice, ABJ; ATP-binding cassette sub-family G member 8, ABCG8; AMP-activated-proteins kinase-α, AMPK-α; AB extracts, ABE; ATP-binding cassette sub-family G member 5, ABCG5; vascular cell adhesion molecule-1 VCAM-1; Apolipoprotein B, ApoB; Area under curve, AUC; Bilberry, BB; BB juice, BBJ; BB extracts, BBE; Black currant, BCT; BCT juice, BCTJ, BCT extracts, BCTE; Blueberry, BlB; BlB juice BlBJ; BIB extracts, BIBE; body weight, BW; C-reactive high sensitivity protein hsCRP; catalase CAT; Cytochrome P450 Family 7 Subfamily A Member 1, CYP7A1; Chokeberry, CB; cranberry, CrB; Cross-over, CO; cross-sectional, CS; cyanidin-3-O-glucoside, C3G; day, d; Double-blind, DB; Freeze-dried, F/D; High-fat-diet, HFD; low-fat-diet, LFD; glucose transporter 1, GLT; glucagon-like-peptide 1, GLP1; Glutathione peroxidase GPx; glutathione reductase GSH-x; 3-hydroxy-3-methylglutaryl-CoA, HMG-CoA; interferon alpha IFN-α; Intercellular Adhesion Molecule 1, ICAM-1; interleukin, IL; Lingonberry, LB; Low-fat-diet, LFD; Low-Density Lipoprotein (LDL) Receptor (LDL-R); Monocyte Chemotaxtractant Protein 1 (MCP-1); Mulberry, MB; Maqui berry, Mqb; Ox-LDL, nonalcoholic fatty liver disease (NAFLD); oxidized low-density-lipoproteins; oxLDL-C; polyunsaturated fatty acids, PUFA; thiobarbituric acid reactive substances (TBARS); total glyceraldehyde, TG; total cholesterol, TC; Tumor necrosis factor, TNF-α; single-blinded, SB; Superoxide dismutase, SOD; one-time-challenge-study, OTCS; placebo-controlled, PC; Peroxisome proliferator-activated receptor-α, PPARα; phosphoenolpyruvate carboxykinase (Pepck); Raspberry, RB; Randomized controlled trial, RCT; respiratory quotient (RQ), short-chain fatty acids, SCFA; sodium glucose transporter protein, SGLT; Sterol regulatory element-binding protein, SREBP-1c; weeks, wk.
Table 2. A comprehensive list of potential health promoting individual anthocyanins and phenolic compounds with their quantities found in berries or berry products.

| Compounds                  | Bilberry (mg/100 g fw) | Blueberry (mg/100 g fw) | Cranberry (mg/100 g dw) | Raspberry (mg/100 g fw) | Mulberry (mg/100 g fw) | Lingonberry (mg/100 g DE) | Blackberry (mg/g DE) | Strawberry (mg/100 g fw) | Goji Berry (mg/100 g dw) | Acai Berry (mg/100 g dw) | Black chokeberry (mg/100 g dw) | Black currant (mg/100 g fw) | Maqui berry (mg/100 g fw) |
|----------------------------|------------------------|-------------------------|-------------------------|-------------------------|------------------------|-------------------------|------------------------|-------------------------|------------------------|------------------------|-----------------------------|----------------------------|--------------------------|
| Cyanidin                   | 62,125                  | 242                     | 135.19                  | 106.6                   | 49.1                   | 593                     |                       |                        |                        |                        |                             |                      |                         |
| Delphinidin                | 105.0                   | 294                     | 120.3                   | 234                     | 230                    | 177.4                   |                       |                        |                        |                        |                             |                      |                         |
| Quercetin                  | 1.5-8                   | 0.07 *                  | 104                     | 0.3-10.04               | 0.09-0.34              | 39.02                   | 37.400                 |                        |                        |                        |                             |                      |                         |
| Myricetin                  | 1.5-8                   | 0.07 *                  | 104                     | 0.3-10.04               | 0.09-0.34              | 39.02                   | 37.400                 |                        |                        |                        |                             |                      |                         |
| p-Coumaric acid            | 1.9                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| m-Coumaric acid            | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Sinapic acid               | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Gallic acid                | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Ascorbic acid              | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Ferulic acid               | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Chlorogenic acid           | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Protocatechuic acid        | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| 5-O-Caffeoylquinic Acid    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                        |                        |                             |                      |                         |
| 1,3-di-O- Caffeoylquinic Acid | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                        |                        |                             |                      |                         |
| Caffeic acid               | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Protocatechuic acid        | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Ellagic acid               | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Vanillic acid              | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Trans-cinnamic acid        | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| O-Hydroxycinnamic acid     | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| p-Hydroxybenzoic acid      | 3.0                     | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Resveratrol                | 1.1-2.1                 | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Epigallocatechin           | 1.1-2.1                 | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| (+/-)-Catechins             | 1.1-2.1                 | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| (+/-)Epicatechin           | 1.1-2.1                 | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Gallic acid gallate        | 1.1-2.1                 | 0.25                    | 67.03-2792.6            | 0.3-4.2                 | 2.64                   | 0.07-0.22               |                        |                        |                        |                        |                             |                      |                         |
| Nutrients 2020, 12, 2538 |
|-------------------------|
| Epigallocatechin         |
| galate                  |
| 1.9                     |
| 4.5-8.4                 |
| 5.65                    |
| Delphinidin 3-          |
| galactoside             |
| 167.1                   |
| 23.4                    |
| Delphinidin 3-          |
| glucoside               |
| 169.1                   |
| 15.4                    |
| 26.8-29.40              |
| 839 ***                 |
| Cyanidin 3-galactoside  |
| 122.6                   |
| 4.2                     |
| 105-2407                |
| Cyanidin 3-glucoside    |
| 130.4                   |
| 2.6                     |
| 998.74                  |
| 327 **                  |
| Petunidin 3-            |
| galactoside             |
| 50                      |
| 11.7                    |
| 103 ***                 |
| Petunidin 3-glucoside   |
| 110.6                   |
| 3.5                     |
| 215-1148                |
| Peonidin 3-galactoside  |
| 101.9                   |
| 12.4                    |
| 10.02-15.25             |
| Peonidin 3-             |
| arabinoside             |
| 13.3                    |
| 1.8                     |
| 2.8                     |
| Peonidin 3-             |
| arabinoside             |
| 23.9                    |
| 9.3                     |
| Malvidin 3-galactoside  |
| 56.7                    |
| 2.1                     |
| 2.04-3.62               |
| Malvidin 3-             |
| arabinoside             |
| 27.5                    |
| 34.9                    |
| 193                     |
| Peonidin 3-             |
| arabinoside             |
| 4.5                     |
| 1                      |
| 4.5                     |
| 4.5                     |
| Malvidin 3-             |
| arabinoside             |
| 67.7                    |
| 31.2                    |
| 9.49-10.57              |
| Malvidin 3-             |
| arabinoside             |
| 12.8                    |
| 34.7                    |
| Quercetin-3-            |
| galactoside             |
| -                      |
| 70.4                    |
| Up to 1.25              |
| Quercetin-3-            |
| α-arabinopyranoside     |
| -                      |
| 34.4                    |
| Quercetin-3-            |
| rhamnoside              |
| -                      |
| 41.6                    |
| Kaempferol-3-           |
| glucoside               |
| -                      |
| 5.6                     |
| 5.12-17.67              |
| Myricetin-3-            |
| α-arabinofuranoside     |
| -                      |
| 37.5                    |
| Quercetin 3-O-          |
| glucuronide             |
| -                      |
| 717.57                  |
| 9.4-39                  |
| Quercetin pentoside     |
| -                      |
| 252                     |
| Cyanidin 3-O-           |
| sophoroside             |
| -                      |
| 43.27-800.3             |
| Cyanidin 3-O-           |
| rutinoside              |
| -                      |
| 5.49-104.58             |
| 433.98                  |
| Pelargonidin 3-         |
| glucoside               |
| -                      |
| 0.14                    |
| Quercetin 3-O-          |
| rutinoside              |
| -                      |
| 192-398                 |
| Quercetin 3-O-          |
| galactoside             |
| -                      |
| 0.2-345                 |
| Quercetin 3-O-          |
| glucoside               |
| -                      |
| 72.4-345.7              |

| Nutrients 2020, 12, 2538 |
|-------------------------|

22 of 66
| Nutrients 2020, 12, 2538 | 23 of 66 |

| Kaempferol 3-O-glucoside | - | - | - | - | 35.5–478 | - | - | - | - | 5.96–14.39 | 0.5–1.94 | - | - | - | - |
| Pelargonidin 3-O-rutinoside | - | - | - | - | 17.8–200 | - | - | - | - | - | - | - | - | - | - |
| Delphinidin-O-(pentosyl)hexoside | - | - | - | - | - | - | 0.82–1.88 | - | - | - | - | - | - | - | - |
| Delphinidin-O-rhamnoside | - | - | - | - | - | - | 2.14 | - | - | - | - | - | - | - | - | - |
| Malvidin-O-pentoside | - | - | - | - | - | - | 1.08–2.13 | - | - | - | - | - | - | - | - |
| Malvidin-O-rhamnoside | - | - | - | - | - | - | 0.13–0.63 | - | - | - | - | - | - | - | - |
| Caffeoylisocitrate | - | - | - | - | - | - | 0.35 | - | - | - | - | - | - | - | - |
| Caffeic acid-O-hexoside | - | - | - | - | - | - | 0.4–0.56 | - | - | - | - | - | - | - | - |
| Myricetin-O-hexoside | - | - | - | - | - | - | 0.19–0.29 | - | - | - | - | - | - | - | - |
| Pelargonidin 3-glucoside | - | - | - | - | - | - | 17.82–20.85 | - | - | - | - | - | - | - | - |
| Pelargonidin 3-malonylglucoside | - | - | - | - | - | - | 5.51–8.16 | - | - | - | - | - | - | - | - |
| Pelargonidin 3-glucoside | - | - | - | - | - | - | 114–348 | - | - | - | - | - | - | - | - |
| Pelargonidin 3-rutinoside | - | - | - | - | - | - | 18–62 | - | - | - | - | - | - | - | - |

Note: Atmospheric-pressure chemical ionization, APCI; Diode array detector, DAD; dry extracts, DE; dry weight, dw; Electron spray ionization, ESI; fresh weight, fw; Hexahydroxydiphenoyl, HHDP; High pressure liquid chromatography, HPLC; Liquid chromatography, LC; Lycium barbarum glycoprotein, LbGp; Lycium barbarum polysaccharides, LBP/LBPC/LBPA/LBPF; Mass spectrometry, MS; Nuclear magnetic resonance, NMR; reverse phase, RP; photodiode array detector, PDA; Quadrupole Time-of-Flight Mass Spectrometry, QTOF-MS; Ultra High pressure liquid chromatography, UPLC. * mg compound/mg extract. ** mg/100 g of sample dw. *** nmol/g..
4. Bilberries

Bilberries (BBs, *Vaccinium myrtillus*) are rich in quercetin, anthocyanins, tannins, catechins, vitamins, and pectins [252]. However, the most important classes of compounds considered responsible for the therapeutic role of BB/BB extracts (BBEE) are phenolic acids and anthocyanins. The majority of compounds belonging to these two classes are presented in Table 2. The phenolics of blueberries varied widely and comprised of 0.3% of fresh fruits, which usually ranged from 48 to 304 mg/100 g of fresh fruit. Among the phenolic acids, the most abundant phenolic acids were ascorbic acid, chlorogenic acids, and 3-caffeoylquinic acid followed by caffeic, ferulic, ellagic, and gallic acids. Among the free phenolic acids, chlorogenic acids and ascobic acids are of prime importance with reference to their health promoting activities [39]. Additionally, at least 15 different BB anthocyanins have been identified including the antidiabetic anthocyanin aglycones, which constituted >70% of the total anthocyanin of BB (Table 2) [17,38,39]. BB anthocyanins showed excellent in vitro α-amylase and α-glucosidase inhibitory activities, reducing or preventing intestinal glucose absorption, and redirecting lipoprotein metabolism regulator enzymatic activities [43]. BB anthocyanins also inhibited advanced glycation end-product (AGE) formation, a severe diabetic complication. The main bioactive compounds considered responsible for inhibiting AGE activity were chlorogenic acid, quercetin-3-galactoside, quercetin-3-arabinoside, quercetin-3-glucoside, quercetin glycoside, quercetin-3-rhamnoside, myricetin glycoside 4, myricetin, and procyanidin b2 biomarkers [253]. BB polyphenols regulate hexose transport via GLUT2 and Na-glucose co-transporter 1 (SGLT-1), which assists glucose uptake. In other studies, GLUT2-mediated hexose transport was impeded by BB-derived flavones [48,254]. Cermak et al. [255] also reported that quercetin-3-O-glucoside and quercetin-4-O-glucoside decreased intestinal hexose absorption by inhibiting SGLUT1 in pig jejunum brush-border-membrane vesicles.

In one of the in vivo studies, supplementation with bilberry extract (BBE) reduced fasting blood sugars (FBS), total glyceraldehyde (TG), TC, and LDL-C levels. BB ingestion increased islet of Langerhans size and minimized retinopathy prognosis. BBE ingestion improved insulin sensitivity and hypoglycemia by upregulating AMPK, which upregulated GLUT4, PPAR-α, ACOX, and carnitine palmitoyltransferase-1 and ACPT-1A, which is synonymous to the suppression of glucose production and increased insulin sensitivity [15]. In another crossover study, the lyophilized BB showed an 18% decrease in (incremental rise of) plasma glucose levels in overweight/obese diabetic humans, accompanied by decreased plasma insulin levels [48]. Recently, Alnajjar et al. [49] also reported that BBE anthocyanins reduced plasma glucose, oral glucose tolerance test (OGTT), TC, high-density lipoprotein cholesterol (HDL-C), LDL-C, TG, and inflammatory adipokine [leptin, TNF-α, and high-sensitivity CRP (hs-CRP)] levels, without affecting the plasma Trolox equivalent antioxidant capacity (TEAC). The anti-inflammatory role of BB was also witnessed when BB juice (BBJ) consumption in healthy adults also reduced NF-κB-regulated inflammatory mediator expression (CRP, IL-6, IL-15, and monokine induced by gamma-interferon) and increased plasma levels of quercetin (by 32–51%) and p-coumaric acid [51]. Later on, Kolehmainen et al. [53] examined the anti-inflammatory mechanism associated with BB consumption and reported the regulation of cytoplasmic ribosomal protein expression and the toll-like receptor (TLR) signaling and β-cell receptor signaling pathways, with decreased proinflammatory macrophage and monocyte functional gene expression including C-C chemokine receptor 2 and monocyte-to-macrophage differentiation. Kim et al. [127] also reported that daily BB consumption reduced vascular permeability by reducing vascular endothelial growth factor levels in diabetic rats, in addition to restoring tight junction protein expression including claudin-5, zona occludens-1, and occludin [127].

An accumulated number of evidence has also suggested that BB(E) intake is also helpful in relieving the oxidative stress and oxidative stress-related complications in obese and (pre)-diabetic subjects (Table 1). BBE administration alleviated stress-induced liver damage by decreasing plasma alanine aminotransferase (ALT), malondialdehyde (MDA), and nitric oxide (NO) levels and increasing glutathione (GSH) and vitamin C levels [45]. Capillary albumin filtration (CAF) is an early diabetic complication, associated with neuropathy and hypertension. BB anthocyanins prevented experimentally-induced-CAF, improving vision and retinopathy, and remarkable CAF reductions.
were observed among diabetic patients [46, 48, 256]. The suggested mechanism for inhibiting CAF involves BB anthocyanosides, which reduced aldose reductase activity and acted as strong antioxidants or pro-reductants, inhibiting AMP and guanosine monophosphate phosphodiesterase by scavenging superoxide anions [256]. Albumin retention (AR) was assessed by the isotopic CAF test in STZ-induced diabetic rats after anthocyanoside-rich BBE administration [46], and BBE treatment was found to reduce and maintain reduced AR (14% to 1.3%) and low-frequency/high-frequency (LF/HF) ratio values in diabetic rats, without toxic effects [47]. BB-derived phenols increased the population of beneficial SCOA-producing gut bacteria (Lactobacillus spp. and Bifidobacterium spp.) and reduced bacterial metabolic syndrome biomarker genera including Enterobacteria. The dysbiosis symbolic Firmicutes/Bacteroidetes ratio, IR, and obesity-led-dysbiosis also decreased following BB consumption [49]. BB added to a fermented oatmeal drink caused a high glucose response, with a significantly reduced insulin index (Table 1) [50].

5. Cranberries

Cranberries (CrBs, Vaccinium macrocarpon) have also been intensively investigated for their proclaimed favorable cardiometabolic and dysmetabolic syndrome effects, likely due to phytochemicals such as oligosaccharides, procyanidins, and anthocyanins. A comprehensive list of potential well-known antioxidant, anti-diabetic, and anti-inflammatory compounds found in CrB (products) or CrB extracts (CrBE) used in clinical or non-clinical interventional studies are listed in Table 2 [226, 227, 228]. The purified fractions of procyanidins were more anti-diabetic potent than the anthocyanin and oligosaccharide fractions [257]. With respect to individual compounds, quercetin-3-galactoside, 5-caffeoylquinic acid, and quercetin-3-rhamnoside were the major compounds comprising 75–77% of total flavonols of cranberry whilst 4-caffeoylquinic acid, 3-caffeoylquinic acid, quercetin-3-arabinopyranoside, myricetin3-gallactoside, myricetin, quercetin-3-arabinofuranoside, and quercetin-3-benzoylgalactoside were found in the least amounts. Many authors have initially described the in vitro anti-diabetic/antiglycation activities of cranberry extracts or its products [226, 257]. Barrett et al. [227] isolated ellagitannins and proanthocyanidins and demonstrated their dose-dependent inhibition of α-amylase and glucoamylase activities. CrB powder from stress-adapted portions of cranberry juice (CrB-JSB) showed increased α-amylase and glucoamylase activities compared with CrB powder, and CrB-JSB (200 mg/mL) also showed anti-hypertensive properties by inhibiting the angiotensin I-converting enzyme (ACE-1) activity [228]. Podsedek et al. [258] found that CrB extracts inhibited pancreatic lipase activities more potently than other berries, but digestive enzyme inhibitory activities were less potent. Purified CrB proanthocyanidins and oligosaccharides also reduced the levels of HbAC1 levels from 7.05% to 5.75, 5.55, and 5.45% in the hemoglobin-glucose assay, whereas the recommended HbAC1 value should be below 7%, according to the American Diabetes Association. Reduced glucose-induced AGE formation during middle glycation stages was also observed during the human serum albumin (HSA)-methylglyoxal and HSA-glucose assays [257]. CrB-derived phenolic-rich extracts decreased fluorescent AGE generation by almost 60%, which was more effective than the other berry anti-AGE activities of raspberries, apples, grapes, and strawberries. The CrB anthocyanin and procyanidin fractions also decreased fluorescent AGE generation in an arginine-methylglyoxal model by 53.3 to 56.8% [226]. The CrB oligosaccharide-rich fraction showed concentration-dependent anti-glycation activity, which reduced AGE formations by 53.3 to 56.8%, respectively, almost as strongly as the reference compound [259].

The hypoglycemic, hypo-insulinemic, and hypolipidemic properties of CrB or its byproducts have also been reported in many clinical interventions (Table 1) [5, 56, 57, 58, 59, 60, 260]. Low-calorie dried cranberry (LCDC, 40 g) consumption after HFD reduced hyperglycemic and hyperlipidemic conditions, halted increases in IR/HOMA-IR and inflammatory biomarkers (TNF-α IL-6, IL-2, IL-10, IL-18, malondialdehyde-MDA) in adipose tissue, and lowered plasma lipid oxidation and oxidative stress biomarker levels in the treated group [56]. After testing LCDC, sweetened, dried CrBs (SWDC) consumed by non-insulinemic diabetic patients also reduced plasma glucose levels when compared with white bread (WB) and unsweetened dried CrBs (USCB) [260]. The plasma insulin peak following
SWDC consumption appeared earlier than the insulin peaks for WB or USCb consumption and was significantly lower than those for WB and USCb. Bread consumption induced higher insulin and postprandial glucose responses, which could be diminished by incorporating CrBs [58,59,60,260]. CrB extracts (CrBEs) also halted visceral adiposity and weight gain in HFD-fed C57BL/6J mice, and improved HFD-induced hypercholesterolemia, hypertriglyceridemia, antioxidant defense mechanisms, and hepatic oxidative stress and normalized the NF-κB/Ilk ratio [54]. Long-term CrBE consumption effects were also investigated [55,63], and the addition of CrBE to normal chow delayed age-related basal plasma insulin concentration declines [63]. CrBE supplementation also improved glucose responsiveness and increased insulin concentrations (7.6%) in rats, without significant HOMA-IR changes. CrBEs also induced duodenal homeobox 1 and insulin expression within islets, which enhanced insulin release, suggesting insulinotropic effect of cranberry intervention [55]. CrBEs showed the anti-obesity effect by inducing the LDL receptor expression, resulting in increased hepatic cholesterol uptake and promoted cholesterol binding to bile acids, causing increased fecal cholesterol excretion [57].

CrBJ consumption was also examined in randomized clinical studies (Table 1). Healthy adults who consumed CrB juice (CrBJ) also showed reduced proinflammatory CRP levels [61]. Daily CrBJ supplementation for 60 days increased paraoxonase-1 (PON-1) and apolipoprotein (Apo)A-I expression (dysfunctioning of PON-1 and apoA-I results in glycation in T2DM patients) accompanied by decreased blood glucose and ApoB levels in T2DM patients. CrBJ inhibited GLUT-1-mediated gastric glucose uptake and aldose reductase, α-amylase, and α-glucosidase activities and protected LDL-C against oxidation [60,64,228]. Moreover, both routine-calorie CrBJ (RCCJ) and high-calorie CrBJ (HCCJ) are enriched in hexoses and sugars, which could limit their use by diabetic individuals. Therefore, low-calorie CrBJ (LCCJ) was examined in glycemic and insulinnemic T2DM patients by Wilson et al. [58,59] and Novotny et al. [65]. LCCJ consumption did not affect LDL-C, HDL-C, or TC levels; however, ApoA-I, ApoA-II, ApoB, and TG levels were reduced in the treated group. Individuals with higher baseline TG or HOMA-IR values experienced more pronounced drops in TG and HOMA-IR than others [65]. Serum HbA1c levels were reduced by 11.4% and 6.02% following RCCJ and RCCJ enriched with omega-3 fatty acid consumption. Omega-3 fatty acid-enriched RCCJ also increased HDL-C levels by 21.1% compared with the baseline [63]. Additionally, folic acid consumption combined with LCCJ decreased plasma homocysteine levels and increased adiponectin and folic acid levels without any change in inflammatory biomarker levels (IL-6, IL-10, IL-18, and TNF-α) [66].

In summary, CrB consumption exerted antimetabolic syndromic effects by downregulating GLUT2 and GLUT4 expression and increasing hepatic cholesterol uptake. Diet-induced weight gain and low-grade inflammation were counteracted by the prevention of TG accumulation and strengthened antioxidative defense mechanisms. The other proposed possible mechanisms of action of CrB, or its products, consumption include reduction and inhibition of ACE-I activity and oxidative stress, accompanied by improvements in endothelium-dependent vasodilation. Furthermore, CrB-derived bioactive compounds including quercetin, inhibited microsomal TG transfer protein (MTP), preventing ApoB-containing lipoprotein assembly. Quercetin also lowered proinflammatory CRP expression in a transgenic mouse model and decreased cytokine-induced CRP expression in Hep3B cells and Chang liver cells [68,69,261], which was analogous to weight loss- and polyunsaturated fatty acid (PUFA)-rich Mediterranean diet-induced CRP suppression [65,261]. Additionally, CrB consumption has beneficial effects on the gut microbiome. HFD reduced Bacteroidetes and increased Firmicutes populations in C57Bl/6J mice, which was reversed by CrBEs intake. CrBEs also increased the Akkermansia gut population, which may prevent HFD-induced increases in circulating proinflammatory lipopolysaccharides (LPS) [54].
6. Raspberries

Raspberries (RBs), especially red RBs (Rubus idaeus L.), are rich in fiber and potent therapeutic phytochemicals that have rendered raspberries as a functional food for metabolic syndrome [199]. The phytochemicals of raspberries provide the healthy and protective affects to its consumers by influencing the cell signaling pathways that affect transporters, receptors, cellular events, and gene expression. These health promoting RB phytochemicals belong to ellagitannins and anthocyanins (Table 2) [262]. Among these two classes, RB anthocyanins are major contributors to health promoting bioactivities. The anthocyanins of RB are cyanidin-based, but with dissimilar glycosidic units. The pelargonidin-based anthocyanins are only found in RB and strawberries with a sophorose unit attachment unique to raspberries. Ellagitannins are hydrolyzable tannins that represent another major RB phytochemical group, which are hexahydroxydiphenoyl esters with quinic acid or glucose cores. Glucose cores can attach to galloyl groups, and further arrangements within hexahydroxydiphenoyl molecules yield the ellagic acids. Numerous in vitro studies have described that RB extracts (RBE) reduced lipid oxidation, LDL-oxidation, ROS generation, and DNA damage, associated with upregulated CAT and SOD enzymatic antioxidant activities [73]. Hypoglycemic studies revealed that RBEs inhibited α-amylase, with mixed effects on α-glucosidase, and aglycones and anthocyanin promoted GSIS from pancreatic cells [263,264]

Fresh RB extracts (RBEs) and freeze-dried RB powder have also been employed for in vivo evaluation (Table 1), in which oxidative stress was found to be relieved as decreased protein and lipid oxidation and damage was seen [74,75]. RB freeze-dried powder fed to obese and diabetic mice reduced ROS levels in erythrocytes by 0.87% when compared to the controls, indicating the ROS-neutralizing role of RB powder bioactive constituents during homeostasis. The RB intervention reduced ROS levels by increasing the glutathione peroxidase (GPx)/SOD ratio (2%) and GPx activity (2.13%) when compared to the placebo controls. Upregulated GPx activity also inhibited lipid peroxidation and protected against diabetes by delaying perturbed metabolism development [76]. RB juice (RBJ) given to hypercholesterolemic golden Syrian hamsters reduced plasma LDLC levels and increased hepatic GSHPx and SOD activities by 30% and 25%, respectively [72]. Polyphenol-rich black RBs have also been combined with HCD foods for sustainable postprandial glycemic control, reducing plasma free fatty acid (FFA) and oxidative stress marker levels. RBs, combined with HCD, blunted postprandial insulinemia and ex vivo LDL-oxidation during the postprandial state, hindering glucose uptake (Table 1) [91]. Purified hydrolyzable RB tannin supplementation in rat gastritis models also demonstrated increased endogenous antioxidant defense system components and decreased inflammatory biomarkers and conditions. RB ellagic acid suppressed the specific immunoglobulin antibody response in cytotoxic cells without affecting other immunoglobulin parameters. Reduced lipid peroxidation, neutrophil infiltration, and iNOS overexpression were observed in ex vivo gastritis and Crohn’s disease models [85,86]. A recent study showed that RBE consumption mitigated carcinogenic acrylamide-induced liver toxicity in male Wistar rats. RB treatment increased plasma antioxidants enzyme levels and reduced acrylamide-induced hepatic ALT, aspartate aminotransferase (AST), alkaline phosphatase (ALP), lactate dehydrogenase (LDH), and gamma-glutamyltransferase (γ-GT) activities [265].

Limited human clinical trials have been performed with RBs, but the antidiabetic effects of RBEs and purified compounds have been examined in diabetic rat models (Table 1) [87,88]. Numerous anthocyanin and polyphenolic components have been hypothesized to affect starch digestion, altering postprandial glucose levels [263]. RB anthocyanin also enhanced insulin sensitivity, upregulated adiponectin expression, downregulated inflammatory cytokines, and altered AMPK phosphorylation, which is a T2DM therapeutic target [264]. A clinical trial examined RB intake with a HCD and reported no postprandial insulin and glucose response alterations [92,93]. In another study following HC-bar consumption, RB intake increased postprandial glucose levels, without changing peak glucose concentrations, and diminished postprandial insulinemia [91]. RB effects on IR and the underlying mechanisms in skeletal muscles were studied by Zhao et al. [87]. AMPK inactivation led to skin lipid accumulation and insulin sensitivity loss. This study found that AMPK-α1 is important for AMPK activation, and dietary RB powder inclusion increased insulin sensitivity.
by upregulating cytochrome C protein in AMPK-α1+/− rats [87]. The supplementation of 5% RB with HFD improved insulin sensitivity by increasing IRS-1 phosphorylation at Tyr 612 and increasing the p-Akt/Akt ratio. RB intake also attenuated nod-like receptor pyrin containing 3 (NLPR3) inflammasome activation, which is a major contributor to metabolic syndrome. NLPR3 activation, combined with caspase 1, forms caspase 1p20 and caspase 1p10. Caspase 1p20 activation releases IL-1β and IL-18. RB consumption downregulated NLPR3, caspase 1p20, IL-1β, and IL-18 expression in HFD-fed mice [88]. Recently, Zou et al. [266] also reported that 5% RB powder supplementation with HFD suppressed TNF-α, L-6, IL-1β, and NF-κB p65 expression and increased GLUT4 expression and IRS-1 and Akt phosphorylation. RB powder also increased mitochondrial biogenesis genes (PGC-1α and Nrf1) and mitochondrial abundance markers (cytochrome c, citrate synthase, and cytochrome c oxidase subunit IV) [266].

The health-promoting effect of raspberry supplementation on the glycerophospholipids metabolism is also evident (Table 1). The addition of 10% freeze-dried RB to an isocaloric diet increased plasma HDL-C (1.5%) and insulin sensitivity and decreased abdominal fat (38%), blood TG, cholesterol, ROS (19%), and LDL-C (0.3%). Similarly, RB-derived cyanidin-3-glucoside upregulated GLUT4 expression, without affecting insulin sensitizer adiponectin [89]. Ellagic acid, which is unique to RB, increased insulin secretion and decreased FBS, HbA1c, and glycated urinary albumin levels. RB inclusion in HFD/HCD diminished impaired insulin tolerance and inflammatory cytokines. RB seed flour, combined with a HCD, downregulated the lipogenic gene expression of lipoprotein lipase (LPL), stearoyl CoA desaturase-1 (SCD-1), and diacylglycerol acyltransferases 2 (DGAT2) and gluconeogenesis promoting genes including PEPCK, G6Pase, sterol regulatory element-binding protein 1c (SREBP-1c), and carbohydrate response element-binding protein (ChREBP) (Table 1) [90]. RB ketones also prevented HFD/HCD-induced BW gains, reduced visceral and adipose tissue, reduced hepatic TG contents, and increased norepinephrine-induced lipolysis in white adipocytes, suppressing lipid accumulation by enhancing lipolysis and fatty acid oxidation [267]. RB supplementation in diabetic patients substantially lowered postprandial glucose levels, without affecting plasma insulin levels after a fatty meal challenge [94]. RB consumption also reduced TG levels [71]. Conflicting results regarding RB interventions and effects on metabolic syndrome biomarkers have been reported. Norrato et al. [76] found an insignificant difference in the weight gain between diabetic mice fed with and without RBs. Similarly, Kirakosyan et al. [77] and Norrato et al. [76] reported no RB intervention effects on LDL-C, fasting blood insulin, IκBα, and PPAR-γ levels. Contrasting results may be due to higher baseline weights of the subjects. However, Kirakosyan et al. showed that RB intake reduced glucose metabolisms and insulin signaling mRNA levels including MAP2K1, glycogen synthase (GYS1), hexokinase, IκBβ, phosphatidylinositol-4,5-bisphosphate 3-kinase, mechanistic target of rapamycin (mTOR), Chuk (involved in innate immunity), C-X-C chemokine receptor type 4 (involved in inflammation), LPL, GYS1, MAP2K1 (involved in apoptosis), nicotinamide phosphoribosyltransferase, ApoE, PPAR-γ, and PPAR-α (involved in glucose and lipid dynamics) (Table 1) [77].

RB intake also increased gut Lactobacillus, which is a healthy gut marker, and increased beneficial gut intestinal SCFAs, which are colonic epithelial cell substrates and improve gut health [78]. RB consumption increased SCFA-producing bacterial populations including Bacteroides, Butyricimonas, Ruminococcus, Akkermansia, Clostridium butyricum, Mucispirillum, Oscillibacter, Ruminococcaceae, and Lachnospiraceae, which improved metabolic syndromic conditions during metformin T2DM treatment [268]. Furthermore, RB consumption time- and dose-dependently increased the gut microbial population of Anaerostipes, Ruminococcus, Akkermansia, Coprobcallicus, Allobaculum, Anaerovorax, Dorea, Asaccharbacter, Anaerotruncus, Coprobcallicus, Desulfovibrio, Victivallis, and Mucispirillum, and decreased the microbial population of Acetivibrio, Anaerotruncus, Bifidobacterium, Lactococcus, Prabacteroides, Streptococcus, Turicibacter, and Acetivibrio. Increased beneficial microbial communities as above-mentioned can reduce inflammation, obesity, metabolic syndrome, and dysbiosis [79]. Su et al. [80] reported that RB-derived pelargonidin-3-O-glucoside increased the gut population of Prevotella and improved the Bacteroidetes/Firmicutes ratio. Another more recent report concluded that there was a favorable higher population of Akkermansia muciniphila and Bacteroidetes/Firmicutes...
Nutrients 2020, 12, 2538

ratios in pathogenic free mice fed on black RB powder [81]. Conclusively, RB consumption showed antidiabetic effects, inhibiting glucosidase and amylase activities, strengthening the endogenous antioxidant defense system, reducing inflammatory biomarkers, activating AMPK, GLUT2/GLUT4, IRS-1 phosphorylation, downregulating lipogenesis and gluconeogenesis genes, and increasing epithelial mucus barrier protecting and SCOA-producing bacterial populations (Table 1) [83,84].

7. Mulberries

Mulberries (MBs, Morus alba/Morus rubra) are rich in cyanidin-3-glucoside, cyanidin-3-rutinoside, and pelargonidin-3-glucoside, and other anthocyanins comprising 78% of the MB polyphenolic compounds (Table 2) [269]. These purified anthocyanins from MB showed excellent glucose-lowering properties in HepG2 cells, increasing PPAR-α and AMPK phosphorylation (activation) and the p-mTOR/mTOR ratio (synonymous with the activation of insulin receptors and insulin-like growth factor 1 receptors). During metabolic syndrome, IRS-1 inactivation increases the p-p38/p38 ratio (subfamily of MAPK, which requires inflammatory cytokines for activation) and reduces PGC-1α expression (a regulator of energy homeostasis and mitochondrial biogenesis), which were abolished or reversed with MB anthocyanins treatment [269]. In addition to anthocyanins, polyphenol-rich MB methanolic extracts also showed excellent α-glucosidase inhibitory activities due to quercetin 3-O-rutinoside, chlorogenic acid, and cyanidin 3-O-glucoside [235]. Cyanidin glycosides in MBs also reportedly possess potent anti-α-glucosidase activity, which inhibit the enzyme by affecting α-glucosidase α-helix contents via cyanidin-3-glucoside (C3G) and cyanidin-3-rutinoside (C3R) domain matching [270]. HepG2 cells treated with the five most abundant MB polyphenols including C3G, 1-deoxyxojirimycin, resveratrol, C3R, and oxyresveratrol showed improved glucose consumption and postprandial glucose disposal through increased glucokinase activity [271]. Another study found that 1,5-dicaffeoylquinic and dihydroquercetin acid protected cells against glucotoxicity [29]. MB extracts (MBEs) upregulated PGC-1α (38%) and FOXO1 (40%) (regulator of PEPCK and G6Pase enzymes) and downregulated PEPCK (79%) and G6Pase (37%) expression in IR model cells. MBEs also upregulated AKT2 (crucial for IRS activation and hence increasing insulin sensitivity) and glycogen synthase kinase (GSK)β3 levels, with significantly increased p-AKT/AKT ratios (hence reduced IR) and increased GSK3β phosphorylation and glycogen synthase 2 (GSY2) activation [272].

In in vivo studies, MB polyphenols and polysaccharides reduced ROS levels and enhanced reductant enzymatic activities including GPx, SOD, and CAT while reducing IL-8, TNF-α, COX-2, and IL-6 release in STZ-induced diabetic mice (Table 1) [273]. MB anthocyanins also attenuated HFD-induced decreased hepatic SOD and GPx activities [95]. Yan et al. [269] reported that MB anthocyanins alleviated hypoglycemia by inhibiting ROS generation, promoting AMPK phosphorylation, activating tuberous sclerosis 2, (reducing the mTOR and ACC signaling), reducing p38-MAPK and PGC-1α expression, and increasing mitochondria and matrix metalloprotease (MMP) abundance in diabetic mice (Table 1) [97]. MB wine consumption by diabetic mice also reversed glycemic status, with reduced oxidative stress markers, proteinuria, non-esterified fatty acid contents, and lipid peroxidation and improved antioxidant defense systems [97]. MB-derived and purified cyanidin-3-O-β-D-glucopyranoside intervention also circumvented diabetic cytotoxicity by reducing oxidative stress markers of DNA modification including 8-hydroxy-2-deoxyguanosine and increasing the axonal transport of nerve growth factor [98].

The oral MBE supplementation also improved insulin signaling by decreased GSK3β, and increased GSY2, AKT, increasing p-AKT/AKT ratios in skeletal, hepatic, and adipocytes tissues of diabetic mice [272]. Oral MB fruit intake in diabetic mice also improved insulin sensitivity by upregulating (up to 3%) the IRS-1, p-IRS01/IRS-1, p-AMPK/AMPK, CCAAT-enhancer-binding proteins (C/EBP), sterol regulatory element-binding protein 1 (SREBP-1c), and PGC-1α [269,274]. Ren et al. [99] further reported that MB consumption normalized glucose metabolism by abolishing protein-tyrosine phosphatase 1B expression and activating the phosphoinositide-3-kinase (PI3K)/AKT pathway. MB anthocyanin-induced p38-AMPK-PGC-1α pathway upregulation increased thermogenesis gene activity. Anthocyanin components also downregulated lipogenesis.
genes including hydroxymethylglutaryl coenzyme A reductase (HMG-CA-R), SREBP-1c, and FAS [100] and activated scavenger receptor class B type 1 and ATP-binding cassette transporter (ABCA1), which transfer cholesterol.

MBEs combined with HFD demonstrated excellent anti-obesity and hypolipidemic properties. MBE supplementation reduced BW gains by 41.3% in HFD-fed diabetic male C57BL/6 mice. Serum TG, TC, HDL-C, and LDL-C levels in HFD + MBE-fed mice were lower than those in HFD-fed diabetic mice, but higher than the MBE-fed controls. Liver injury parameters (ALT and AST) were reduced in HFD + MBE-fed mice, with reduced adipose and hepatic liver lipid droplet sizes [101]. MB fruit consumption lowered TG, TC, LDL-C, and FFA levels in other studies (Table 1) [102,103]. MB-derived anthocyanin consumption decreased serum levels of inflammatory markers (IL-6, IL-1α, iNOS, TNF-α, IFN-γ, and NF-κB), thiobarbituric-acid-reactive substances (TBARS) (a lipid oxidation marker), hyperlipidemic markers (TC, glucose, TG, and leptin), insulin, and hepatic AST, ALP, and ALT levels, downregulated FAS, and increased heme oxygenase-1 (HO-1) (a cytoprotective enzyme) and antioxidant enzyme levels in HFD-fed male C57BL/6 mice (Table 1) [95,104]. Aqueous MBEs employed the hypolipidemic and hypoglycemic effects by activating the AMPK, increasing the p-AMPK/AMPK ratio (hence improving mitochondrial biogenesis), and downregulated FAS, acetyl coenzyme A carboxylase (ACC), glycerol-3-phosphate acyltransferase (GPAT), and SREBP-1 [104]. MBEs in HFD-fed male Sprague-Dawley rats prevented non-alcoholic fatty liver disease (NAFLD) by downregulating lipid/cholesterol homeostasis-related genes (FAS, ACC, GPAT, and SREBP-1) and suppressing the lipid oxidation biomarkers MDA and 4-hydroxynonenal [105,106]. Hu et al. [275] demonstrated that MBE increased nuclear factor erythroid-2-related factor 2 (Nrf2) phosphorylation and nuclear translocation, activating the Nrf2/antioxidant response element signaling pathway, which increased quinone oxidoreductase 1, HO-1, and NAD(P)H expression and promoted antioxidant enzymatic activities, thus protecting hepatocytes against palmitic acid-induced lipo-toxicity and oxidative stress.

Gut microbiota regulates dietary energy harvesting, glucose homeostasis, and lipid metabolism, especially in brown adipose tissues (BAT). Mitochondria-rich BAT activation can increase energy expenditure following MB-induced UCP1 upregulation and oxidative phosphorylation downregulation, releasing energy as heat. MB powder consumption reversed HFD-induced gut microbiome changes, increasing the Bacteroidetes/Firmicutes ratio and Bacteroidetes populations (Porphyromonadaceae, Parabacteroid, S24-7, Prevotellaceae, Alloprevotella, Rikenellaceae, Alitest, Rikenella) and decreasing the Proteobacteria (Alphaproteobacteria, Brevundimonas, Devosia, Rhodobacteraceae, Polymorphaobacter, Deltaproteobacteria, Desulfuovibrio, Arsenimonas), and Firmicutes (Clostridia, Lachnospiraceae, Eubacterium, Coprococcus, Ruminococcaceae, Oscillibacter, Ruminoclostridium) populations [107,108]. At the genus-level, MB fruit supplementation promoted SCOA/SCFA-producing and IMBD-restoration-supportive genera Lactobacillus, Bacteroides, Bacteroide, Allobaculum, and Akkermansia growth, and suppressed Corynebacterium, Staphylococcus, Aerococcus, Jeogalicoccus, Facklamia, and Enterococcus growth. Allobaculum and Lactobacillus protect against metabolic syndrome, and both genera increased in diabetic rats after MB intake [108]. Approximately 60 metabolites were identified in MB including flavonols, phenolic acids, flavonoids, lignans, and organic acids (Table 2) [234]. In short, MB fruit consumption upregulated/activated glucose-consumption-related pathways and insulin-sensitivity-related pathways (p-AKT/AKT ratio, glucokinase, PGC-1α, FOXO1, IRS-1, p-IRS-1/IRS-1, p-AMPK/AMPK, C/EBP, and Bacteroidetes/Firmicutes ratio) and downregulated lipogenesis-related pathways (FAS, ACC, GPAT, and SREBP-1) in skeletal, hepatic, and adipocyte tissues.

8. Lingonberries

Lingonberry (LB, Vaccinium vitis-idaea) alleviates metabolic syndrome including frequent urination and fatigue. In in vitro studies, LB extracts (LBEs) increased glucose uptake in C2C12 skeletal muscle cells by modulating AMPK activity [276]. LB polysaccharides inhibited α-glucosidase activity (by 118–136%) more strongly than the referenced acarbose [277]. In in vitro digestibility assays, LB polyphenols (7% w/v) were added to white rice, which significantly reduced glucose
release [278]. Ethanolic LBEs demonstrated antiglycation activity, with AGE inhibition majorly mediated by LB cyanidin-3-galactoside, quercetin-3-galactoside, and (+)-catechin [279]. In J774 macrophages, LBEs significantly inhibited LPS-modulated NO production, without substantial effects on COX-2 or iNOS expression. Proinflammatory cytokine (IL-6, IL-1β, and TNF-α) expression was reduced by TNF-α downregulation, IkB receptor degradation inhibition, and reduced extracellular signal-related kinase 1/2 phosphorylation [280]. However, in RAW 264.7 macrophages and activated 3T3-L1 adipocytes, LBEs mitigated oxidative stress by suppressing COX-2, iNOS, TNF-α, IL-6, MCP-1, and IL-1β expression [281].

In in vivo studies, LB consumption also improved hyperinsulinemic, hyperglycemic, and dyslipidemic conditions (Table 1) [111]. LBE consumption reduced blood glucose levels (17–25%), obesity-induced hepatic steatosis (50–60%), and plasma TG, TC, and LDL-C levels (12–18%) associated with increased GLUT4 expression and AMPK and Akt phosphorylation, increasing glucose metabolism and hepatic fatty acid oxidation [111]. LB juice (LBJ) improved low-grade inflammation and endothelial function by increasing NO availability, which is necessary for the inhibition of adhesion molecules, MCP-1, ACE-1, COX-2, and other pro-inflammatory markers [112]. The LB-rich Okinawan-based Nordic diet improved anthropometric (BW, body mass index (BMI), and waist circumference) and metabolic (HOMA-IR, IR, FBS, TG, CRP, TC, and HDL-C) parameters [119]. Linderborg et al. [120] demonstrated that LB powder consumption compensated for additional glucose and lipid consumption. LB intake prevented HFD-induced BW gains in C57BL/6J mice. LB supplementation reduced FBS, fasting insulin, and HOMA-IR levels (Table 1) [113,114]. Hepatic lipid accumulation and liver function parameters (ALT, TG, and cholesterol) decreased after LB supplementation, more strongly than other berries [113,114]. In a recent hyperlipidic and hypercaloric meals challenge study, the LB supplementation halted increased cholesterolemia and decreased the glycemic response, CRP, and postprandial endotoxia [121]. In an atherosclerosis ApoE−/− mouse model, whole LB consumption upregulated bile acid synthesis gene Cyp7a1, increased the cecal propionic-acid-producing bacteria proportions, and decreased triglyceridemia and atherosclerosis [115]. The insulinenic and glycemic response following oat bread consumption was also checked. The LB polysaccharide and fiber consumption, following bread consumption, reduced glucose and CRP responses [122]. Whole LB and LB nectar intake reduced postprandial glucose and insulin levels after 35 g sucrose intake, and insulin levels increased more rapidly following LB than after glucose intake. Postprandial glucose levels were also reduced following LBJ consumption. Insulin and FFA changes after LBJ consumption were similar to those observed after whole fruit consumption (Table 1) [113,114,123].

Urinary metabolomics revealed that a LBJ-containing diet increased 4-hydroxyhippuric acid and hippuric acid excretion, whereas 4-deoxythreonic acid, 3-hydroxybutanoic acid, dimethylamine, creatinine, and citric acid excretion reduced, likely due to high polyphenolic compound and benzoic acid contents in LBJ (Table 2) [124,282]. Plasma lipidomics data showed that LB consumption increased health-promoting lyso-phosphatidylethanolamines, (LPE) (16:0), lysophosphatidylcholine (LPC) (20:5), (16:1), and (22:5), and phosphatidylcholines (PC) (33:2), (32:2), (35:6), (34:4), (36:6), and (36:5), whereas obesity and diabetes symbolic sphingomyelins (SM) (34:1), (33:1), (40:3), and (38:2) were reduced. Quinate levels also increased, and plasma alanine and glucose levels decreased significantly [116]. LBE and powder supplementation of HFD downregulated the expression levels of macrophage marker endothelial growth factor-like module containing mucin-like, hormone receptor-like 1 (EMR1), and LPS-sensing TLR4 (member of the toll-like receptor family activation of which results in signaling the NF-κB pathway and inflammatory cytokine production) and upregulated tight junction-associated occluding (an integral membrane protein whose modulation is associated with cellular proliferation, differentiation, signal transduction, and migration) and proglucagon (a precursor of glucagon from α-pancreatic cells). The HFD-fed control microbiome showed the upregulation of the ATP-binding cassette (ABC) transporter, cell motility, membrane transporter, bacterial chemotaxis, bacterial motility, the two-component system, flagellar assembly, transcription, and signal transduction genes, compared with the LB-treated group [283]. LB consumption enriched genes associated with lipid metabolism, nutrient transport, energy,
nutrients, and amino acids (Table 1) [113,114,117]. At the phyla level, LB supplementation affected the diversity and population of Firmicutes, Bacteroidetes, Proteobacteria, and Verrucomicrobia. The relative abundance of Bacteroidetes increased, and the relative abundance of Firmicutes decreased significantly, reducing the obesity and diabetes symbolic Firmicutes/Bacteroidetes ratio [113,114,117]. At the genus level, HFD increased Firmicutes genera populations including Lachnospiraceae, Oscillospira, and Ruminococcus. The abundance of Bacteroidetes increased following LB supplementation, due to unknown members of the S24-7 family. LB supplementation increased Parabacteroides, Odoribacter, and Akkermansia populations. The principal component analysis confirmed LB extract-induced gut microbial profile variations. HFD increased the population density of the genera Oscillospira and Ruminococcus and the Lachnospiraceae family, microbes associated with diabetes pathogenesis progression [284], which was prevented by LB fruit/powder/extract consumption [285]. Akkermansia population increases were associated with the abundance of Akkermansia muciniphila species, which are known beneficial gut microbacteria that counteract HFD-induced adipose tissue inflammation, endotoxemia, BW gain, and IR in C57BL/6 mice [286]. Liquid chromatography (LC)-tandem mass spectrometry (MS/MS)-based LB fingerprinting identified several bioactive compounds responsible for antioxidative, antidiabetic, and anti-inflammatory properties. These bioactive compounds primarily belong to anthocyanidins, flavonols, glycosides, catechins, and different conjugates of ferulic and caffeoyl acid (Table 2). Depending on aglycon weight, cyaniding-containing compounds were the major bioactive compounds followed by proanthocyanidins, which represent phenolic compounds in LB [236,237].

9. Blackberries

Blackberries (Rubus grandifolius L and Rubus fruticosus L.) are consumed fresh or as juices, jams, and liqueurs. Blackberries are enriched in health-promoting compounds (Table 2) belonging to flavanols, flavanones, flavonols (kaempferol and quercetin glycosides), anthocyanins, hydroxycinnamic acids, and caffeic acid conjugates. The high-performance liquid chromatography (HPLC)-electrospray ionization (ESI)-mass spectrometry (MS)-based Rubus grandifolius L. metabolic profiling revealed 50 phytochemicals including anthocyanins, hydroxycinnamic acids, flavonols, flavanones, and ellagitannins (Table 2) [131,238,239]. These blackberry-derived compounds offered an antidiabetic and anti-obesity role by inhibiting digestive enzymes (α- and β-glucosidase, aldose reductase, lipase, and α-amylase) and exhibiting anti-glycation abilities. The blackberry α-glucosidase and α-amylase inhibitory activity was superior to the reference compounds, Acarbose and 1-Deoxynojirimycin (1-DNJ) [287]. Anthocyanins are considered to be the primary mediator of blackberry extract anti-digestive activities, and glycosides are the primary inhibitors of α-glucosidase activity. The interaction between glycosides and enzymes is considered to be competitive, suggesting that glycosides bind to enzymatic active sites [288]. Cytidine glycosides from leaf and fruit R. grandifolius extracts reduced aldose reductase activity, which is responsible for AGE accumulation in diabetic patients via dicarbonyl activity [289]. The recorded anti-glycation activity of BB fruit extracts was IC50 = 1.87 mg/mL, and ellagitannins and flavonols were the most prominent anti-glycation agents [131,238,239]. HepG2 cells incubated with gut microbial-fermented blackberry metabolites (GMBB) and gastrointestinal-digested BB slurry (GIDBB) showed improved glucose uptake. Increased HepG2 uptake also increased glycogen synthesis. GIDBB and GMBB also maintained the desired cellular redox status by neutralizing ROS and restoring the mitochondrial membrane potential. GIDBB and GMBB supplementation restored glutathione levels, strengthening the oxidative defense system [290].

In in vivo studies, blackberry-derived purified anthocyanin-enriched and ellagitannin-enriched fractions decreased lipid peroxidation markers (TBARS and MDA) and increased hepatic and brain antioxidant enzyme activities (CAT, GSH, SOD, and GPx) [125]. Similarly, blackberry extract consumption attenuated the HFD-induced effects in an obesity-prone mouse model and prevented the increase in metabolic and lipidemic parameters, while reinforcing endogenous and exogenous antioxidant enzyme systems (Table 1) [126]. LPL activity, plasma glucose, insulin, and acyl-carnitines were also upregulated after blackberry consumption. Antioxidative enzyme system reinforcement
correlated with the anti-inflammatory and anti-dyslipidemia potential of blackberry extracts [127]. The glycemic and lipidemic-controlling mechanisms of blackberry extracts were mediated through the downregulation of lipogenesis factors (FAS, SCD-1, microsomal triglyceride transfer protein, diglycerides acyltransferase, and adipose triacylglyceride lipase), energy coupling/uncoupling proteins (UCP-1, UCP-2, and UCP-3), pro-inflammatory cytokines (PPAR-α, Nrf2, IL-6, and TNF-α), and fatty acid β-oxidation genes (CPT-1α and ACOX-1) (Table 1) [127], which were maintained by long-term and chronic blackberry extract consumption. Additionally, the increases in total monounsaturated fatty acid contents of adipocytes, plasma brain-derived neurotrophic factor levels, and pro-inflammatory leptin levels in HFD-fed controls were counteracted by blackberry extract consumption [128]. Human clinical trials were also run, in which healthy human subjects were given BB fruits in addition to HFD, resulting in reduced fat accumulation and increased fat oxidation. Blackberry consumption lowered postprandial glucose and lipid levels by activating AMPK and BAdTs. [291]. Pulpy blackberry juice consumption by dyslipidemic patients decreased ApoB and hs-CRP, increased ApoA-1 and HDL-C, and left other lipid parameters unaffected [134]. In healthy subjects, blackberry juice increased exogenous and endogenous antioxidant enzymes. Cyanidin, ascorbate, total ellagic acid, urate, and R-tocopherol contributed to increased plasma and urine antioxidant capacities [125,133]. Daily blackberry consumption reduced dyslipidemia and insulinemic parameters in diabetic and obese adults [132]. Blackberry polyphenolic compounds inhibit digestive enzyme activities, physically interacting with hexose absorption transporters and modulating transporter expression at the genomic level [292]. Blackberry compounds may also modulate peripheral glucose use, damaged pancreatic cell regeneration, and enhance blood glucose withdrawal by increasing insulin sensitivity (Table 1) [109,129].

Blackberry juice was also examined in STZ-induced-diabetic male Sprague-Dawley and hamster rats (Table 1) [129]. Blackberry juice significantly reduced food and water intake, reducing the BWs of both control and diabetic rats [129]. Blackberry nectar supplementation of a cholesterolemic diet reduced hyperlipidemic parameters and hepatic lipid peroxidation [181]. Blackberry juice consumption effectively reduced triacylglycerols (~43.5%), glucose (~48.6%), and cholesterol (~28.6%) levels without side effects. Blackberry juice consumption limited lipid peroxidation in the plasma (~7.5%) and kidneys (~19.5%). Similarly, alcohol-free fermented blackberry juice (AFBBJ) was used to supplement HFD in obese C57BL/6j mice [270], which significantly reduced fat-mass gain and FBS and decreased plasma TG, TC, LDL-C, and HOMA-IR levels, while increasing β-cell function (HOMA-β) [22]. Liver function tests revealed no change in ALT, but AST increased in AFBBJ-treated mice. Genomic sequencing approaches revealed pancreatic gene upregulation, responsible for amino acid and glucose metabolism and insulin secretion regulation [22].

The intestinal bioavailability of blackberry polyphenols and resulting impact on gut microflora have also been recently investigated. The low-absorption and cecal accumulation of BB polyphenols were the main reasons for positive health effects. The cecal microbial fermentation of blackberry polyphenols generates antidiabetic and antioxidative blackberry metabolites including C3G, 2,4,6-trihydroxybenzoic acid, coumarin, and caffeic acid. The increased cecal glycoside concentration and secondary metabolites improved glucose consumption (Table 1) [290]. The increased cecal SCFA concentration suggested an increase in SCFA-producing bacteria; however, the relative abundance of different bacterial groups was not reported [130]. Blackberry treatment altered the gut microfloral composition by increasing cecal Bacteriodetes over Firmicutes. Lactobacillus johnsonii was abundant in both blackberry-treated and control groups, whereas Lachnospiraceae dominated the blackberry group, promoting glycoside metabolism. However, Clostridiales, Enterococcus faecalis, and Bifidobacterium pseudolongum were more dominant in the control groups [131].
10. Strawberries

Strawberry (Fragaria × ananassa) consumption has been associated with decreased risk and occurrence of metabolic syndrome, cancer, diabetes, chronic inflammation, and hypertension. The credit of these health-promoting activities goes to its rich phytochemical contents (Table 2). Many studies analytically analyzed the crude and fractionated phytochemical contents of strawberry and found strawberry rich in antioxidative, anti-obesity, antiglycation, anti-inflammatory, and antidiabetic compounds from flavanols, flavonols, anthocyanins, hydroxyxynamic acid derivatives, hydroxybenzoic acid derivatives, ellagic acid and ellagic acid glycosides, and ellagittannins (Table 2). The most surplus glucose-lowering acid moieties were malonic and p-coumaric acid and the most identified flavonols of strawberry were derivatives of kaempferol and quercetin glycosides. The red-coloration-granting and anti-oxidative anthocyanins of strawberries were mostly the derivatives of pelargonidin and cyanidin [240]. The hydrolysis of ellagittannins gave rise to the most important antidiabetic phytochemical called ellagic acid, which comprised more than 50% of the total polyphenolic components of strawberry. The level of ellagic acid is about 3–10 times higher in the strawberry than other berries, fruits, and nuts. It is one of the constituents due to which strawberry can regarded as a functional food [293]. In in vitro studies, strawberry ethanolic extracts inhibited pancreatic lipase activity more strongly than reference orlistat. Aqueous and ethanolic strawberry extracts inhibited adipocyte cell division and inhibited inflammatory mediator (β-hexosaminidase and histamine) release by 61.8 to 80% [294]. Strawberry polyphenolic compounds interact with glucose transporters such as SGLT1 and GLUT2 and attenuate glucose uptake due to polyphenol compound competition for transporter active sites [295]. HPLC-diode array detector (DAD)-MS analysis and statistical correlations showed the contribution of pelargonidin-3-O-glucoside to glucose uptake inhibition. Strawberry extracts effectively inhibited uptake and transport of glucose up to 5% in HepG2 cultures [295]. Da Silva Pinto et al. [296] showed that the strawberry extract α-glucosidase inhibitory activity was superior to the α-amylase inhibitory activity. Strawberry-derived ellagitannin consumption (>50 mg/mL) sufficiently inhibited ACE activity [296]. Methanolic strawberry extracts activated p-AMPK/AMPK expression in HepG2 cells, resulting in fatty acid and cholesterol regulatory gene inactivation and phosphorylation including HMG-CoA-R and ACC. Activated p-AMPK/AMPK expression increased LDL receptor expression including PGC-1α and sirtuin 1 (a NAD-dependent deacetylase that inhibit hepatic lipogenesis, stimulating FA β-oxidation, and maintaining cholesterol and bile acid levels) in HepG2 cells [297].

Numerous in vivo studies have also cited the health promoting activities of strawberry or its byproducts in animal models and human clinical trials. The intake of aqueous, alcoholic, and hydro-alcoholic strawberry extracts improved the serum glucose level, liver function (decreased serum glutamic pyruvic transaminase, serum glutamic oxaloacetic transaminase, alkaline phosphatase), lipid profile (decreased LDL-C, LDL-C/HDL-C, and LDL-C/TC ratio), and lipid oxidation markers (decreased MDA and CAT) [136,137]. Genes associated with glucose, cholesterol, and lipid metabolism [FAS, ACC, CPT-1A, malonyl-CoA, acyltransferase, ACC-α (ACACA), and acyl-CoA synthetase long-chain family member 1] were also downregulated by strawberry treatment [135]. Paquette et al. [143] used the hyperinsulinemic-euglycemic clamp methodology to examine improved insulin sensitivity after strawberry extract consumption, but did not detect improvements in fasting insulin and glucose concentrations. In animal studies, HFD supplementation with strawberry prevented weight gain without influencing food and water intake. Strawberry beverage consumption protected against postprandial lipemia by reducing TG (14%), TC (5%), and LDL-C levels (5%) in hyperlipidemic patients following HFD [144]. Sugar-rich strawberry jam consumption also attenuated glycemic index and postprandial glucose level increases in diabetic human subjects [145,298]. Strawberry jam consumption showed favorable lipid and sugar metabolism results, even compared with low-sugar strawberry jam [146]. Strawberry consumption with HCD also controlled postprandial glucose levels, affected glucose and insulin responses, and GLP-1 expression. Regular strawberry beverage and juice consumption decreased blood pressure, TC, and the TC/HDL-C ratio in diabetic patients. T2DM and CVD risk factors were also ameliorated (Table 1) [66,114,123,147]. Strawberry extracts reduced IL-6 and plasminogen activator inhibitor 1...
(PAI-1) (a risk factor for atherosclerosis) levels in obese individuals after HFD/HCD, without influencing TNF-α, CRP, platelet aggregation, or fasting insulin and glucose levels [148]. In another similar study, the postprandial insulin level and inflammatory response (hs-CRP and IL-6) were reduced with increased plasma pelargonidin sulfate and pelargonidin-3-O-glucosidein levels after strawberry powder consumption with high-carbohydrate, moderate-fat meals [149]. In another recent study, strawberry-blueberry powder, consumed with a HFD/HCD, reduced BW gains (12.7%), visceral fat mass (18%), retropitoneal and subcutaneous white adipose tissues (up to 10.45–16.5%), postprandial insulin and glucose levels, IR, and inflammatory markers (MCP-1, TNF-α, IL6, CRP, and PPAR-α), in male Wistar rats and C57BB/6j mice (Table 1) [19,138]. Strawberry-blueberry powder exerted anti-adipogenic effects by regulating lipid metabolizing genes including PPAR-α and C/EBPα. Inflammatory and lipogenesis-related gene expression were reduced including TNF-α, IL6, and C/EBPα, adipogenesis-driver transcription factors (PPAR-γ), adiponectin, adipocyte fatty acid-binding protein, SREBF1, leptin, SCD-1, and FAS [138]. In another dose-response checking study, the intake of strawberry against the Western-type-meal reduced the oxidized low-density lipoproteins and post-meal insulin demand in insulin resistant patients [150].

Oxidative stress is a leading cause of metabolic syndrome and diabetes. Strawberry powder supplementation in an isoenergetic diet containing the oxidative-inducing antibiotic drug doxorubicin reversed doxorubicin-induced decreases in the antioxidants retinol and α-tocopherol and upregulated liver antioxidant enzymes including GPx, CAT, GSH, SOD, and GST (Table 1). Plasma hepatic stress biomarker levels including protein carbonyls and hydroperoxide were reduced by strawberry intake [139,152]. Strawberry-based foods containing carbohydrate, fat, and lipids increased total antioxidant levels (1.26 to 1.45 mmol/l) of the subjects while decreasing HbA1C (from 7.00 to 6.72%) levels. The plasma hs-CRP and MDA levels also decreased from 3.36 to 2.76 mmol/mL and 3.36 to 2.76 nmol/mL, respectively [153]. Strawberry powder intake prevented HFD- and stress-induced decreases in γ-aminobutyric acid levels and reduced oxidative stress and lipid oxidation markers, in male Wistar rats [140]. Fresh strawberry consumption reduced linseed oil-induced DNA damage and plasma oxidative marker levels and increased the plasma antioxidant status of pigs [299].

Strawberry intake effects on gut microbial ecology in diabetic subjects increased phylogenetic species richness (α-diversity) and global microbial composition (β-diversity) variations at the genus and operational taxonomic unit levels. Proteobacteria, Actinobacteria, and Verrucomicrobia were significantly altered after the strawberry intervention. Strawberry intake significantly increased the abundance of beneficial Bacteroides and Actinobacteria and decreased Akkermansia, Verrucomicrobia, Dehalobacterium, and Dorea (Firmicutes). At the genus level, the abundance of SCOA-producing Lactobacillus and “prebiotic-effect-giving” Bifidobacterium increased, whereas Dehalobacterium, Dorea, SMB53, and Turicibacter remained unaltered [141]. Additionally, a specific relationship between ingested flavonoids and microbial community patterns was identified [151]. Dietary flavanol and flavanone intake were positively associated with Eggerthela lenta. Flavonols and flavanol monomer intake was positively associated with Adlcreutzia equilicafiens (involved in phytochemical degradation) and inversely associated with Flavonifractor plautii (Gram-negative poorly understood) populations [151]. Whole strawberry powder intake increased the α-diversity of colonic inflammatory CD-1 mice, increasing Bifidobacterium and Lactobacillus and reducing pro-inflammatory Akkermansia, Dorea, and Bilophila [142]. The polyphenolic compounds that affected gut microbiota compositions in strawberry fruit extracts were flavonols, flavonol anthocyanins, hydroxycinnamic acid derivatives, hydroxybenzoic acid derivatives, ellagic acid, ellagic acid glycosides, and ellagitannins (Table 2) [240,241].

11. Goji Berries

Goji berry (GB, Lycium Barbarum) is a functional food and alternative therapeutic tool for T2DM treatment [155]. The major GB therapeutic phytochemicals include polysaccharides (5–8%), carotenoids (0.03–0.5%), and phenolic compounds (traces). The compounds belonging to these classes have been listed in Table 2 [123,242]. The GB is considered the best source of diapalmitin zeaxanthin carotenoids. These carotenoids showed effective protection against diabetic-induced-retinopathy
The in vitro hypoglycemic tests showed the inhibitory capability of GB carotenoids was 9.6 to 82.6% and 5.7 to 15.3% for α-glucosidase and α-amylase enzymes, respectively [242]. In GB polyphenolic compounds, phenolic acids (24.7%) and flavonoids (75.3%) are major phytochemical classes. The major therapeutic flavonoids in GB are squercetin-3-O-rutinoside (from 7.1 to 232.7 mg/kg) and quercetin-3-O-hexoside (from 169.1 to 1107.7 mg/kg) whereas phenolic compounds include caffeoylquinic acid (0.34 µg/g), caffeic acid (3.73 µg/g), p-coumaric acid (6.06 µg/g), chlorogenic acid (12.4 µg/g), kaempferol-3-O-rutinoside (11.3 µg/g), quercetin-diglucoside (66.0 µg/g), and rutin (42.0 µg/g) [242]. As GB polysaccharides (GBPS) are major contributors of health-endowing activities and have been vastly investigated, this section will primarily focus on GBPS. GBPS are considered to be therapeutic in alternative medicine with immunomodulation, antioxidant, neuroprotection, anti-tumor, antidiabetic, radioprotection, anti-osteoporosis, hepatoprotection, and anti-fatigue activities. The GBPS biological activities depend on their molecular weight, chemical structure, and chain conformation [154,301]. The GBPS are among a few plant-based bioactive compounds that have shown simultaneous hypoglycemic and hypolipidemic properties. Due to hypoglycemic and anti-hyperlipidemic properties, GBPS may be a potent T2DM inhibitor, delaying disease prognosis, even after disease development. Antidiabetic assays showed impressive lipid and glucose reducing effects [155,302]. Acidic GBPS treatment in rat insulinoma cells decreased oxidative stress biomarkers and increased antioxidant enzyme systems. GBPS treatment of IR alloxan-treated-HepG2 cells protected against oxidative stress and improved cell survival and proliferation [302]. Similarly, the GBPS was further checked for possibly hampering glucose uptake in the gut and intestine. The GBPS intensively reduced glucose absorption in a dose-dependent manner by competing for intestinal absorption [303]. Rat insulinoma cells incubated with GBPS rescued damaged pancreatic cells, improved the survival rate, and encouraged insulin secretion. The IR cell model was supplemented with purified GBPS, which upregulated glucose consumption. GBPS was easily translocated and transported across the Caco-2 intestinal cell membrane through the SGLT-1 transporter, producing a hypoglycemic effect. Therefore, GBPS is a plant-based bioactive compound that shows simultaneous hypoglycemic and hypolipidemic properties [303]. Purified GBPS fractions showed dose-dependent hypoglycemic activities, resulting in increased glucose uptake [156,303]. Besides GBPS, GB carotenoids have also shown antidiabetic and α-glucosidase and α-amylase enzyme inhibitory activities [242].

The hypolipidemic effects of GB intake have been studied by in vivo approaches (Table 1), but human clinical trials for GB have been limited, with most studies performed using small sample sizes in China. GB consumption effectively reduced serum lipid peroxide species in diabetic patients. Reductions in waist circumference, TG, transaminase, and TC levels were reported in metabolic syndrome patients following routine GB intake. Lipid profile improvements were accompanied by increased GSH and CAT enzymatic activities [167]. GB anthocyanins reduced BW gain (17.4 to 38.7%) by increasing fecal fatty acid contents and downregulating IL-6, TNF-α, IFN-γ, NF-κB, and iNOS gene expression [157]. GBPS decoction treatment of alloxan-induced, diabetic, obese rabbits effectively reduced blood glucose levels. GBPS substantially decreased serum TG (~4.27%), TC (~3.5%), LDL-C levels, and increased HDL-C serum levels (0.78) [154]. The hypoglycemic and hypolipidemic effects of GBPS were later confirmed by the works of Zhao et al. [158]. Supplementation of HFD with GBPS decreased HOMA-IR, fasting and postprandial insulin and glucose levels, serum TG, TC, and LDL-C levels, and weight gain [158].

The oxidative stress relieving effect of GBPS was also checked (Table 1). The effect of GBPS treatment on the kidneys of STZ-induced diabetic rats increased kidney antioxidant enzymes including CAT, SOD, GPx, GST, and GSH [170]. The supplementation of GB in the form of GB milkshakes increased plasma zeaxanthin and antioxidant levels by 57 and 26%, respectively. GB juice (GBJ) also increased GSH peroxidase (GSH-Px) and SOD by 9.87% and 8.7%, respectively and decreased MDA levels by 5.95% [166]. GBJ intake also protected against glaucoma, which was confirmed in retinal ganglion cells, and disrupted intraocular pressure [159]. GBPS administration to C57BL/6 mice reversed oxidative stress, dyslipidemia, and diabetic changes. GBPS administration downregulated nitrotyrosine and MDA expression and increased antioxidant enzymes such as CAT,
**Nutrients 2020, 12, 2538**

GPx, and Cu/Zn SOD. GBPS intake also diminished pro-inflammatory biomarkers including TNF-α, IL-1β, iNOS, and COX-2. Following pro-inflammatory marker reduction, liver injury biomarkers, called chemokines, were also reduced. The liver regeneration process was also observed following GB intake, enhancing liver regeneration biomarkers [168,304].

With respect to hypoglycemic effect specifically (Table 1), Zhao et al. [160] confirmed the antidiabetic characteristics of GBPS, which increased GLUT-4 expression in the skeletal muscle plasma membrane. Purified GBPS in pancreatic cells increased glucose uptake and metabolism, insulin secretion, and proliferation. The enhanced glucose metabolism mechanism was associated with increased hepatic hexokinase and pyruvate kinase expression/activity (Table 1) [26,161]. GBPS may block the ATP-sensitive K+ channel, activate glycogen synthetase and insulin-like growth factor, enhance peripheral glucose utilization, or inhibit glucagon releasing factors in pancreatic α-cells [197]. In a recent single meal challenge study, increased glucose and lipid consumption were observed in GB-treated patients, associated with increased respiratory quotients, oxygen usage, and carbon dioxide release. However, no single-dose effects on substrate oxidation and postprandial-energy-expenditure were reported [169]. Du et al. [162] compared GBPS with metformin and reported similar normalization effects on blood glucose and insulin levels. This study also reported reduced IL-2, IL-6, TNF-α, intercellular adhesion molecule-1 (ICAM-1), MCP-1, and blood urea/nitrogen levels, inhibited albuminuria, and reversed histopathological alterations. GBPS treatment in HFD/HCDFed rats also demonstrated hypoglycemic and hypolipidemic effects [115]. Ni et al. [163] examined the potential neuroprotective effects of aqueous GB extracts. Retinal apoptosis causes photoreceptor degradation and diabetic retinopathy (DN), and GB carotenoid supplementation in rats hampered caspase-2-induced apoptosis, protecting photoreceptors [163]. Prolonged or chronic hyperglycemia downregulates luteolin and zeaxanthin-metabolizing gene expression, causing retinopathy. GB carotenoids protected against diabetes-induced retinopathy. GB supplementation upregulated carotenoid metabolism genes and retina biogenesis in STZ-induced diabetic rats [300]. GB also contains taurine, a non-essential amino acid, and GB-derived taurine enhanced PPAR-γ activity and elevated cAMP levels, hampering the prognosis of DN with reversal of epithelial barrier impairments [300].

GBPS, polyphenol, and carotenoid effects on the gut microbiome were also studied (Table 1). Fermentation and simulated digestion experiments revealed that GBPS was digested and degraded only in the distal gut, releasing monosaccharides and promoting beneficial SCOA-producing bacterial growth. Monosaccharides with side chains are more susceptible to degradation than monosaccharides with linked backbones. GBPS greatly increased SCFA-producing gut microbiota and increased Bacteroidetes (including Prevotella and Bacteroides) and Actinobacteria (containing Collinsella and Bifidobacterium) populations, whereas Megamonas and Megasphaera (Firmicutes) populations were decreased. Furthermore, SCOA/SCFA-producing, prebiotic-effect-giving, proteolytic microflora such as Bacteroides, Phascolarctobacterium, Bifidobacterium, Prevotella, Clostridium XIVb, Oscillibacter Collinsella, and Lactococcus were prominent following GBPS treatment [305]. In another study, dietary GB supplementation also increased health-promoting secondary metabolite and SCOA-producing Actinobacteria, Lachnospiraceae, Clostridium XIVb, Sporobacter, Pseudoflavonifractor, Butyricicoccus, Anaerotruncus, Anaerospirabacter, and Ruminococcaceae populations without affecting Akkemansia, Mucispirillum, Bacteroides, and Desulfovibrio. Butyryl-Coenzyme A CoA transferase is an important butyrate gene, and GBPS supplementation increased its expression in butyrate-producing bacteria such as the Clostridium cluster XIVa group including Lachnospiraceae, Faecalibacterium prausnitzii, and Ruminococcaceae [164]. The GBPS prebiotic effects increased the populations of Firmicutes, Akkemansia, Proteobacteria, Lactobacillus, and Prevotellaceae [165].
12. Acai Berries

Acai berry (AB, *Euterpe oleracea*) is native to South America and has high phytochemical contents. The dominant antidiabetic phenolic acid constituents in AB include ferulic acid, anthocyanin-3-glycosides, *p*-hydroxybenzoic acid, epicatechin, protocatechuic acid, gallic acid, ellagic acid, catechin, *p*-coumaric acid, vanillic acid, and galactomannans (Table 2) [246]. Anthocyanin and flavonoids are prominent therapeutic polyphenols including C3G and C3R [244,245]. AB juice (ABJ) is richer in polyphenols and flavonoids than other berry juices, resulting in increased antioxidant capacities [200]. In in vitro studies, the isotonic ABJ pancreatic lipase inhibitory activity was significantly positively correlated with anthocyanin contents. Isotonic ABJ also reduced adipogenesis and lipid accumulation in 3T3-L1 adipocytes and inhibited α-glucosidase activity [306]. Isotonic ABJ also inhibited Cu-mediated LDL oxidation and oxidized or acetylated LDL uptake. AB puree also showed antiglycation activities at a concentration 0.1 mg/mL, which was 89% stronger than the control [171]. Polyphenols in ABJ affect adipogenesis, preventing obesity, weight gain, inflammation, and diabetes [307].

In in vivo studies, AB fruit proved to be a very useful therapeutic agent for circumventing oxidative stress, and controlling dyslipidemic and metabolic syndrome conditions (Table 1). The supplementation of AB fruit effectively prevented protein oxidation as increased protein sulfhydryl groups were observed, with decreased protein oxidation biomarker carbonyl proteins. A single AB pulp dose enhanced plasma antioxidant capacity 7-fold 3 h after its consumption. Plasma anthocyanins reached maximum levels 2.2 h after AB pulp consumption [179,180]. In another in vivo study, AB pulp supplementation in oxidatively damaged mutant *Drosophila melanogaster*, in combination with HFD, reversed HFD-induced oxidative stress damage and prolonged the lifespan expectancy by 22% [172,308]. AB supplementation with exercise improved hepatic oxidation status by reducing inflammatory MCP-1 expression, SOD activity, redox-sensitive signaling pathway activation, ROS generation, and ROS stress [173]. To elucidate the antidiabetic and antioxidative molecular mechanism of AB, AB-mediated transcript-level changes were examined in 12 genes associated with JNK, nutrient sensing, and insulin-like signaling pathways [309]. PEPCK genes, involved in glyceroneogenesis and gluconeogenesis, were reduced in the AB pulp group. Cholesterolemic diet consumption decreased lethal/essential or life gene (lefl2) expression, which was reversed by AB fruit consumption. Two JNK targets, metallothionein A, and glutathione S transferase D1, which have antioxidant activities, were upregulated after AB consumption without affecting the remaining JNK downstream target genes (Ferritin 1 heavy chain homolog, Ice, Heat shock protein 68, and Puckered). Moreover, AB ingestion promoted longevity by intensifying stress response pathway activity and suppressing PEPCK genes [172,309]. Treatment with AB seed extracts also reduced blood pressure, the hypertension biomarker renin, and DN biomarker levels (creatinine, urea, creatin, and albumin). Diabetes onset leads to oxidative stress and hypertension, decreasing the number of glomeruli per area per kidney, a major DN marker. AB seed extracts reduced kidney volume expansion and prevented a decrease in the number of glomeruli per area per kidney [174]. AB seed extracts substantially reduced renal injury (resulting in reduced urea and creatine excretion), hampering renal fibrosis progression. The diabetes-induced glomerular filtration barrier injury markers, podocin and nephrin, decreased in diabetic male Wistar rats, whereas AB seed extract treatment restored these levels. AB seed extract treatment also reduced renal proinflammatory cytokines and oxidative stress biomarkers, reinforcing the anti-oxidative defense system [174]. The effects of exercise and AB seed-rendered extract consumption in STZ and HFD-induced diabetic rats reduced HbA1C, glycemia, serum insulin, HOMA-IR, serum TG, TC, LDL-C, and HDL-C levels [176]. Insulin signaling components (insulin receptors, pAKT, and AKT) in skeletal muscles were upregulated following AB seed extract consumption and exercise [176]. Reduced adiponectin levels are observed in T2DM, associated with deregulated sugar and lipid metabolism, and AB seed extracts reversed this effect. AB seeds induced increased GLUT-4 expression and glucose uptake due to AMPK activation [176] and increased GLP-1 and incretin levels with reduced leptin and inflammatory cytokine expression, which were not observed in HFD-fed rats treated with exercise alone. Increased GLP-1 and incretin expression promotes insulin secretion, suppressing gastric
emptying, and glucagon synthase [176,310]. The same research group then used the AB seed extracts to check the anti-obesity features in the C57BL/6 mice strain fed on HFD. HFD supplemented with the AB seed extract prevented weight gain in mice [311]. Adiponectin levels, which are responsible for lipid metabolism, decreased in HFD-fed mice and were restored by AB seed extract supplementation. AB seed extracts increased glucose and lipid metabolizing protein expression including pAMPK/AMPK, pACC/ACC, HMG-CoA, and various transporters including ATP-binding cassette sub-family G member 5-ABCG5 and ATP-binding cassette sub-family G member 8-ABCG8, while reducing SREBP-1c expression. Similarly, protein and lipid oxidation products including carbonyl proteins and MDA were reduced by strengthening the anti-oxidative enzyme system [311].

Regarding glucose-lowering effect, recently, the human AB fruit consumption with normal meals decreased FBS and mean plasma insulin levels after one month. Plasma TG, TC, and LDL-C levels, and the LDL-C/HDL-C ratio also decreased, with increased plasma HDL-C levels [181]. The AB consumption with HFD enhanced fecal cholesterol contents, with no influence on low-grade-inflammation biomarkers [113]. Freeze-dried AB fruit pulp reversed the HFD-induced alterations in PEPCK expression [312]. Aqueous ethanolic AB extracts restored mitochondrial complex I function by modulating NADH:ubiquinone oxidoreductase core unit 7 and 8 expression. NLRP3 (a component of inflammasome) and caspase 1/caspase 3/caspase 8 (Interleukin-1 converting enzyme family, which initiates inflammatory response) were downregulated in oxidative-agent-treated macrophages [313]. AB supplementation also interfered with hepatic cholesterolemic metabolism. AB attenuated the high-cholesterol diet effects by reducing weight gain, TC and LDL-C levels, and key regulatory gene expression associated with the cholesterol biosynthesis pathway including HMG CoA-R, EBP-2, ApoB100, LDL-R, ABCG8, and CYP7A1 [175]. Intensive feeding with freeze-dried AB pulp attenuated HFD-induced hepatic steatosis by improving IR, adiponectin expression, adiponectin receptor 2, SREBP-1c, PPAR-α, and its target gene, CPT. Fat accumulating gene expression including UCP-2 and fatty acid translocase were reduced by AB treatment [179]. Both lipid accumulation and oxidation were reduced in zebrafish fed with a high-cholesterol diet, and reduced serum TC, LDL-C, and MDA levels were observed in AB-treated zebrafish [171]. Aside from lipid oxidation inhibition, the AB intake also prevented amino acid oxidation after HCD, reducing protein carbonyls and sulfhydryl groups, which are important protein damage biomarkers. Reduced arylesterase and PON activities and reduced hepatic ALT, AST, and ALP levels demonstrated improved hepatic operation [175]. AB powder also improved anti-inflammatory mechanisms after HFD by improving glucose intolerance and reducing IL-6 and TNF-α concentrations in epididymal adipose tissue [312].

A comprehensive study examining AB intake on the gut microflora is currently lacking. Simulated digestion studies examining AB polyphenols inhibited the growth of symbiotic and saccharolytic Bacteroides, Prevotella, and Clostridium histolyticum. AB polyphenols showed favorable effects on the intestinal SCFA bacteria population including LAB [178]. Guerzotto et al. [177] noted increased intestinal populations of obesity-protecting bacteria (i.e., Bifidobacterium spp., Eubacterium rectale–Clostridium cocoideis group, Bacteroides spp – Prevotella group, and FOS-Rafitilde P95). However, AB polyphenols showed no considerable effects on Enterococcus spp and C. histolyticum [177].

Conclusively, AB exerted antidiabetic, anti-obesity, antioxidative, and anti-inflammatory actions by reducing the expression of PPAR-γ and its modulators (C/EBP-β, C/EBP-δ, and other C/EBP family members, Kruppel-like factor, and SREBP1C) Moreover, decreased expression level of transcriptomic factors such as C/EBPβ (-0.41%), C/EBPα (-0.66%), Kruppel like factor (-0.83%), and SREBP1C (-0.24%) were also seen [125,133]. AB also reduced the expression levels of lipogenic genes FAS (-0.5%), aP2 (-0.7%), LPL (-0.7%), and FATP1 (-0.55%). Low-grade-inflammation biomarkers including leptin and total PAI decreased with increasing anti-inflammatory and anti-adipogenic adiponectin levels [170,172,309,314]. The expression levels of the pro-inflammatory factors NF-κB, TNF-α, MCP-1 (-0.81%), IL-6 (-0.48%), IL-8 (-0.05%), IL-1β (-0.03%), and INF-β (-0.49%) were also reduced. TNF-α activates NF-κB and interleukins (IL-2 and IL-6), which was prevented by AB polyphenols [170,313,314].
13. Chokeberries

Chokeberries (black chokeberry (BCB), *Aronia melanocarpa*, red chokeberry (RCB), *Aronia arbutifolia*) can be consumed as whole fruit, jam, wine, juice, syrup, tea, soft spreads, chili starters, salsa, beer, extracts, gummies, ice cream, and tinctures. CB consumption was used to treat colds in America and to treat hyperglycemia, metabolic syndrome, and hypertension in Europe and Russia. In in vitro bioassays, CB extract (CBE) showed significant α-glucosidase inhibitory activity compared with the referenced antidiabetic drug acarbose. Purified anthocyanins (cyanidin 3-galactoside, cyanidin 3-arabinoside, cyanidin 3-glucoside, and cyanidin 3-xyloside) were the strongest antidiabetic compounds compared with isolated dimeric and trimeric procyanidins. BCB juice (BCB) also inhibited α-glucosidase, dipeptidyl peptidase (DPP) IV, and ACE activities by 75, 35, and 95% in a dose-dependent manner, respectively [182]. BCB fermentation and digestion increase polyphenol bioaccessibility. Fermented and digested *Aronia* kefir showed stronger α-glucosidase (IC₅₀ = 152.53 ± 15.24 mg kefir/mL) and pancreatic α-amylase inhibitory (IC₅₀ = 146.52 ± 5.37 mg kefir/mL) activities than non-fermented *Aronia* (IC₅₀ = 365.16 ± 370 48.84 mg and 196.21 ± 5.50 mg, respectively) [315]. BCB relieved oxidative stress in βTC3 cells by restoring the anti-oxidative enzyme pool and insulin secretion, as comprehensively explained in Figure 2 [316]. The oxidative-stress-induced reduction in insulin secretion was restored by the BCB extract (BCBE) treatment under basal glucose conditions [316]. BCBE treatment of pancreatic cells nullified cytokine (IL-1β and IFN-γ)-induced effects and decreased oxidative stress production [183]. BCBE pretreatment (0.001, 0.01, 0.1, or 1 mg/mL) of diabetic hepatic cells line RINm5F reduced cytokine-induced-oxidative stress from 19.3–0.39 μM to 14.9–0.35 μM [183]. Similarly, BCBE pretreatment of HAECS nullified the TNF-α-induced ICAM-1 and VCAM-1 expression by 35 and 45%, respectively, in a dose-dependent manner. BCBEs also prevented NF-κB p65 phosphorylation, which activates the pro-inflammatory transcription factor NF-κB [317,318].

![Figure 2. Schematic presentation of chokeberry anthocyanin-induced insulin secretion and antioxidant enzyme pathways in pancreatic β-cells under high-glucose-induced stress conditions.](image-url)

Glucose is transported across the cell membrane via glucose transporter (i.e., GLUT-2), followed by glycolysis and pyruvate production. Afterward, pyruvate is used for the generation of ATP in mitochondria. Here, in connection with the electron transport chain, radicals, like superoxide anion (O₂⁻), are also produced and simultaneously neutralized by the enzymatic antioxidant SOD. SOD converts the O₂⁻ into harmless O₂ and another radical H₂O₂. In addition to H₂O₂ diffusion through the cell membrane, H₂O₂ is also scavenged by CAT and GPx resulting in water and oxygen production. Chokeberry-derived anthocyanins strengthen this inherent enzymatic antioxidant system (i.e., SOD, CAT, and GPx), which can more actively neutralize the radicals generated during glucose metabolism. H₂O₂-stimulated reduction of GSH is also ameliorated by chokeberry anthocyanins. Chokeberry anthocyanins also replenish the pool of insulin by increasing the insulin gene expression.
Proinsulin, a precursor of insulin, folded in the endoplasmic reticulum, is transported to the Golgi apparatus. Chokeberry anthocyanins can also influence the opening of the voltage-gated Ca\(^{2+}\) channels, leading to an increased fusion of insulin granules with the cell membrane (Source: Rugina et al. [316]).

Addressing the anti-inflammatory potential of CB, in in vivo clinical studies, Kardum et al. [195,196] administered CBJ to patients with pharmacologically incurable grade I hypertension and high blood pressure, resulting in decreased systolic/diastolic blood pressure, with a stronger effect associated with long-term consumption. CBEs also reduced systolic/diastolic blood pressure [197], particularly in congenital heart disease patients [198]. Following hypertension, inflammation is another diabetes complication and numerous studies have cited the anti-inflammatory potential of BCB or its juice consumption. Increased PPAR-\(\gamma\)2 expression was attenuated by BCBEs, reducing downstream lipid metabolizing PPAR-\(\gamma\)2 target expression such as PGE receptor and LPL, decreasing intracellular lipid droplet accumulation [184]. Regular BCBJ consumption improved chronic inflammatory conditions, lowering IFN-\(\gamma\) and TNF-\(\alpha\) levels [195,196,198]. The immunomodulatory effects of BCB intake have also been discussed in the literature in STZ-induced male Wistar rats. DM causes immune imbalances because damaged pancreatic cells trigger macrophage and T lymphocyte infiltration, which lesion \(\beta\)-cells. BCB consumption by STZ-induced male Wistar rats reduced fibrinogen, TNF-\(\alpha\), and IFN-\(\gamma\) levels, which returned to their normal values 72 h post-administration of BCB [199].

Regarding hypoglycemic response, BCBJ consumption also modulated circulating lipid levels including TG, TC, and LDL-C in mild hypertensive patients (Table 1) [185,198]. BCBJ consumption also reduced serum TG, TC, and LDL-C levels in hypercholesterolemic healthy subjects [128]. Long-term BCB consumption was recommended for desirable hypoglycemic and hypolipidemic effects [128,185,198]. Valcheva-Kuzmanova et al. [186] demonstrated up to 39% reduced postprandial serum TG levels in STZ-induced diabetic rats after BCBJ consumption and reported encouraging results for both diabetic and healthy rats. However, Lipińska and Jóźwik [187] showed pronounced hypolipidemic effects only in diabetic Polish Merino lambs including significantly decreased serum LDL-C and increased HDL-C levels, without significant effects on serum TC levels. In addition to preventing increased plasma glucose, homocysteine, and fibrinogen levels, reduced serum lipid levels (TG, TC, and LDL-C) were observed in STZ-induced diabetic rats [201]. Hepatic steatosis and NAFLD were prevented by BCB treatment in HFD-fed diabetic C57BL/6N mice. Daily BCBE administration prevented increased body, liver, and epididymis weights [188]. Several possible mechanisms have been proposed in the literature referring to the lipid-lowering property of BCB consumption. The BCB hypoglycemic effect may be associated with increased cyanidin-induced lipid metabolism, reduced catechin-induced cholesterol absorption, and the flavonoid-influenced downregulation of cholesterol synthesis enzymes including HMG-CA-R, cholesterol acyltransferase, and acyl-CoA [185,188].

The anti-oxidative, anti-obesity, and anti-diabetic potential of BCB was checked in the various diabetic model mice (Table 1), where BCB increased serum insulin secretion with reduced pro-inflammatory cytokine expression (MAPKs, NF-\(\kappa\)B, COX-2, and iNOS) in a dose-dependent manner [183]. Jurgoński et al. [189] fed BCB to high-fructose-diet-fed STZ-induced diabetic rats and showed increased maltase and sucrase activity, and decreased lactase production in the small intestinal mucosal membrane. Daily BCBJ consumption lowered postprandial glucose levels after OGTT, regardless of gender, and reduced ACE, \(\alpha\)-glucosidase, and DPP IV activities in a dose-dependent manner [182]. Valcheva-Kuzmanova et al. [186] showed lower postprandial glucose levels (up to 44%) in STZ-induced diabetic rats after BCBJ consumption, and Lipińska and Jóźwik [187] demonstrated a pronounced FBS decrease in BCB-treated Polish merino lambs. Postprandial OGTT results for BCB-treated mice decreased, with improved intraperitoneal ITT results [185]. Similarly, consumption by STZ-induced diabetic mice reduced serum TBARS levels and mitigated lipid peroxidation (by 29–50%) and kidney hypertrophy [190]. Following CCl\(_4\) administration, the decreased concentration of CAT, GPx, and GR were increased by 117%, 56% and 44%, respectively, after the intake of BCBJ. Protein carbonyls, protein oxidation biomarkers, decreased by 22% after
BCBJ consumption in male Wistar rats [190]. BCBJ consumption by the KK-Ay and C57BL/6jmsSlc mice reduced BW, white adipose tissue weight, α-glucosidase and DPP IV activity, and blood TG levels. Mesenteric, epididymal, subcutaneous, and retroperitoneal white adipose tissue weights were reduced by 26%, 27%, 48%, and 38% compared with those in control animals [39]. Bhaswant et al. [191] administered BCBJ to male Wister HFD- and HCD-fed rats and observed reduced BW gain and feed conversion efficiency. Total body fat mass, BMI, abdominal fat (epididymal, omental fat pads, and retroperitoneal), and visceral adiposity index reductions were more pronounced in Wistar rats fed with BCBJ than in those fed with biofunctional purple maize flour. BCBJ consumption also reduced liver injury biomarkers (ALP, AST, and ALT), although these levels remained within the normal range [191]. In another study, male Wistar rats were fed high-fructose diets containing CBCE, resulting in increased plasma HDL-C and adiponectin levels [192]. IRS-1/2 and PI3K regulatory subunit protein expression increased by 2.3-, 1.8-, and 1.5-times, respectively, along with inhibiting the phosphatase and tensin homolog (Pten) (~0.61%) expression. The expression level of glucose uptake, transportation (GLUT1 and GLUT4) and gluconeogenesis (GYS) was uplifted by 1.5 times compared to high-fructose fed control rats. BCBJ consumption inhibited lipogenesis and lipid accumulation by reducing fatty acid-binding protein, FAS, and LPL (lipogenesis protein) by 0.6–0.7%. Improved glucose and lipid metabolism and increased glucose and lipid regulatory metabolizing protein expression (adiponectin and PPAR-γ) were also observed [192]. Cyanidine 3, 5-diglucoside was identified as a DPP IV inhibitor. DPP IV cleaves incertins including GLP-1 and glucose-dependent-insulino-tropic polypeptide at their N-terminal regions, resulting in decreased insulin secretion [182,186]. Cyanidin glycosides including 3-galactoside, 3-glucoside, cyanidin 3-O-β-glucoside3-arabinoside, and 3-xylloside enhance glucose uptake and GLUT4 translocation. Diabetes-associated hyperlipidemic complications were improved by regulating the FOXO1-mediated adipose TG lipase transcription [185].

BCBJ contains high levels of anthocyanins (1958.18 mg/100 g FW), proanthocyanidins (522–1002 mg/100 g FW), and hydroxycinnamic acids (187.9 mg/100 g FW) including chlorogenic acid and neochlorogenic acid [212,247]. Cyanidine-3-O-glucoside, cyanidin-3-O-galactoside, cyanidine-3-O-xylloside, and cyanidine-3-O-arabinoside are the primary antidiabetic and anti-oxidative anthocyanin compounds in BCBJ (Table 2). No studies have examined the CB consumption effects on gut microbiota in diabetic/obese individuals, although CB consumption has been examined in healthy individuals [202]. Chronic BC capsule treatment influenced the intestinal diversity of health promoting and SCOA-producing Anaerostipes, Bifidobacterium, Faecalibacterium, and Clostridium genera. CBE capsules increased the relative abundance of Anaerostipes, whereas whole CB capsules increased Bacteroides and Clostridium XIV populations. Correlation analysis between gut microbial genera and plasma polyphenolic contents revealed that Prevotella, Dialister, Desulfovibrio, and Bifidobacteria were responsible for the increased levels of nine, eight, seven, and six health promoting plasma CB metabolites, respectively, including derivatives of benzoic acid, hippuric acid, phenylacetic acid, cinnamic acid, caffeic acid, flavonols, (iso)ferulic acid, benzaldehydes, and pyrogallol [202].

14. Black Currants

Black currant (BCT,Ribes nigrum L.) is cultivated primarily in Europe, New Zealand, and Australia. BCT is a rich source of anthocyanins that represent 95% of polyphenolic compounds, with the remaining 5% including other minor polyphenol classes. Delphinidin-3-rutinoside (D3R) is the major BCT antidiabetic anthocyanin compound that improves glucose tolerance. In BCT nectar, cyanidine and delphinidin rutinosides are the dominating anthocyanins, followed by glucoside compounds [93,319]. A full list of other therapeutic BCT compounds are presented in Table 2. GLP-1 and AMPK are the primary BCT polyphenolic compound targets. BCT extract (BCTE) consumption increased GLP-1 secretion. GLP-1, an incretin, promotes pancreatic β-cell division and glucose-dependent insulin release [212,213,289]. BCTEs contain approximately 70% anthocyanins (especially rutinosides and glucosides of delphinidin and cyanidin) and are considered to be effective α-glucosidase inhibitors [289]. Apple and BCT juice (BCTJ) treatment in human Caco-2 cells reduced
sodium-independent and total glucose uptake by 46 and 51%, respectively. In oocytes, apple and BCTJ-derived phloretin and phlorizin effectively reduced glucose uptake by 58 and 85%, respectively [213]. The BCT polysaccharide BCP-I also showed remarkable antiglycation activities due to its inhibitory effects on Amadori products [320]. BCT powder incorporation into high-glycemic-indexed food decreased glucose release and increased antioxidant capacities [321].

In addition to in vitro studies, glucose and lipid lowering effect of BC extracts or its screened anthocyanins have also been investigated enormously in various in vivo studies (Table 1). The intake of major BCT anthocyanin consumption, in combination with intraperitoneal glucose administration, prevented increased serum glucose concentrations with the simultaneous increase in serum insulin levels [203]. Improved hyperglycemia and hypoinsulinemia are caused by the GLP-1 activation-induced increase in insulin secretion. BCT powder, administered for six days before OGGT, improved postprandial plasma insulin and glucose levels in healthy human subjects [214]. BCTE consumed with a normal diet by KK-Ay mice induced hypoglycemia and modulated basal GLP-1 concentrations without affecting plasma insulin levels, food intake, or BW [204]. Proglucagon cleaving agent proprotein convertase subtilisin/Kexin type 1, which processes proglucagon into GLP-1, increased. BCTE treatments also increased AMPK phosphorylation in skeletal muscles, upregulating insulin-independent glucose uptake pathways by increasing downstream target expression including GLUT-4 and the translocating plasma membrane [204]. Previously, Esposito et al. [205] also conducted an anti-diabetic study using 1% BCT powder, which decreased rat BWs, irrespective of dietary fat contents. Microbiological fecal analyses showed increased fecal anthocyanin contents, especially in lean animals. These results suggested that gut microflora more actively transform polyphenolic metabolites in lean animals rather than in obese animals. BCT supplementation reversed the postprandial glucose levels associated with HFD; however, the postprandial glucose level continued to rise due to gut microbiota disruption. Similarly, BCT improved HFD-induced insulin, but the gut microflora disruption increased IR. These results signified the importance of gut microflora during the BCT polyphenol metabolization and biotransformation [205]. The supplementation of 0.1% BCTE in HFD reduced retroperitoneal and epididymal adipose fat. BCTE hypolipidemic characteristics were verified by upregulated lipogenic/lipid metabolizing genes in adipocytes including UCP-2, UCP-3, mitochondrial transcription factor A (TFAM), PPARY, SREBP-1c, PPARα, and SCD-1, and fatty acid oxidation genes including CPT-1α and 1β [206]. Repressed inflammatory marker expression in macrophages has also been reported. Reduced IKKε (an enzyme complex that is involved in propagating the cellular response to inflammation) and TANK-binding kinase 1 (a member of IKK subfamily, which activates in response to lipopolysaccharides) expression was observed in the BCT-treated group, compared with upregulation in the HFD group [206,212]. BCTJ/nectar waste extract (pomace) was much richer in anthocyanins than in BCT pulp. Phytochemically, BCT pomace extracts are rich in D3G, D3R, cyanidin-3-rutinoside, glycosides, and flavonol aglycones. HFD supplemented with BCT pomace extracts did not affect food intake or BW. Fat in the diet increases small intestinal digesta viscosity, whereas BCT pomace polyphenolic extracts made this digesta more acidic [207]. The polyphenolic-rich BCTE also reduced cecal tissue mass and increased ammonia contents. HFD reduced bacterial glycolytic enzyme activities such as α- and β-galactosidases and α- and β-glucosidases, which were restored by BCT pomace extract. BCT supplementation reduced β-glucuronidase activity, which is associated with reduced pressure on the intestinal detoxification mechanism [208]. BCT supplementation reduced the cecal putrefactive SFCA concentration, regardless of diet [207,208]. BCTE consumption increased mean fat oxidation during prolonged cycling exercise by endurance-trained females with reduced mean carbohydrate oxidation [215]. However, the opposite outcome was observed when BCTJ was consumed before exercise, without significant effects on blood lactate, glucose, and MDA levels [216].

In addition to HFD, the high-fructose-diet or HCD were also involved in the hyperglycemic, hyperlipidemic, and metabolic syndrome conditions. BCTE administration with high-fructose-diet prevented increases in liver weight, BW, and epididymal fat pad weight. OGTT results improved, with decreased p-AMPK and IRS-1 levels in the BCTE-treated group. BCTE supplementation also decreased high-fructose-diet-induced hyperglycemic marker expression and reduced atherosclerosis
risk by diminishing ICAM-1, VCAM-1, E-selectin, endothelin, and eNOS expression levels in aortic tissues [209]. Consumption of an anthocyanin-rich sugar-free BCT drink with a normal-carbohydrate diet delayed the glycemic and insulinemic response with reduced incretin and GLP-1 expression [212,213]. The consumption of BB, BCT, CrB, and strawberries restricted post-meal blood insulin and glucose fluctuations induced by HFD/HCD. LB combined with BCT (whole or nectar) ameliorated postprandial insulinemic and glycemic control and response [62,123,267]. The irreversible hydrolysis of sucrose into fructose and glucose under high temperature and low pH conditions produces invertase sugars. BCT nectar, sweetened with invertase sugars, reduced postprandial blood glucose levels and the maximal blood glucose level by 33 and 87%, respectively. The nectar x time interaction also revealed lower insulin secretion at 15 and 30 min of post-nectar-consumption and expulsion of insulin from the baseline was cut by 13% compared to the reference [62,123,267].

Regarding oxidative stress and diabetes-related complications, ample amounts of evidence have suggested that anthocyanins from BC exert anti-hypertensive, anti-inflammatory, anti-fibrotic, and anti-hepatic steatosis effects by limiting lipogenesis and gluconeogenesis (Table 2) [217]. BCT-derived purified extracts administered to hepatic steatosis model C57BL/6J mice did not prevent BW loss, but serum ALT and AST levels increased. BCT anthocyanin supplementation decreased hepatic TG and TC accumulation [304]. Histological analysis showed that microvascular steatosis, inflammatory cell infiltration, and hepatocyte ballooning were reduced by (up to 50%) BCT anthocyanins. Hepatic stellate cells produce collagen during fibrogenesis. Reduced α-smooth muscle actin and upregulated carbamoyl phosphate synthase 1 suggest hepatic stellate cell inhibition, inhibiting fibrosis and non-alcoholic hepatic steatosis. BCTE treatment increased mitochondrial biogenesis and decreased the AMPK/βAMPK ratio and pivotal mitochondrial biogenesis regulators including PGC-1α and β, Nrf-1 and -2, and TFAM. Mitochondrial fatty acid β-oxidation occurs due to mitochondrial oxidative phosphorylation, which was reversed through effects on PPAR-α, CPT-1, and medium-chain acyl CoA dehydrogenase expression [62,93,123,217,304,319].

15. Maqui Berries

Maqui berries (MB) (Aristotelia chilensis) have recently gained attention due to their high content of polyphenolic compounds. The stated phytochemical composition of MB was 138 ± 0.4 mg/100 g fresh weight with 35% relative abundance of delphinidin [218]. Di Lorenzo et al. [219] analyzed the MqB composition (Table 2). MqB is rich in anthocyanins including 84% diglycosylated and 16% monoglycosylated anthocyanins [251,322,323]. The in vitro sugar hydrolyzing enzymes inhibitory activities of MB extracts were reported by Rubiliar and his colleagues. Rubiliar et al. [324] reported α-amylase and α-glucosidase inhibitory activities, resulting in decreased postprandial glucose levels and improved glucose tolerance [324]. Crude and purified MqB extracts (MqBEs) reduced MDA production and minimized oxidative damage [250]. An isotonic soft drink containing lyophilized MqB, acai, and blackthorn berry powders [152] demonstrated pancreatic lipase and α-glucosidase inhibitory activities, which were superior to the control, acai-, and blackthorn-based beverages. Likewise, the in vitro anti-diabetic assay showed the inhibition of α-glucosidase activity by 90% compared to the lemon juice control (80%), whilst the recorded inhibitory α-glucosidase activity of tested commercial isotonic drinks was around 50% [306]. The MqBE anti-diabetic and anti-lipidemic potentials were further examined in RAW264.7 mouse monocytes and 3T3-L1 mouse pre-adipocytes [220,325]. MBEs reduced adipocyte formation by promoting MMP-2 and MMP-9 (endopeptidases). GST treatment decreased GSH, SOD, and CAT expression, which was reversed by MqBE treatment in macrophages. LPS treatment increased IL-6, MCP-1, TNF-α, and galectin-3 with decreased adiponectin expression, which was countered and reversed by MqBEs in macrophages [220]. Furthermore, a dose of 100 and 180 μM MqB delphinidin inhibited sodium palmitate-induced-TG-accumulation by 50 and 59%, respectively, in HepG2 cells [222].

With respect to in vivo antidiabetic and anti-obesity potential of MqB (Table 1), Rojo et al. [220] fed C57BL/6J mice anthocyanin-rich MqBEs, which significantly decreased plasma glucose levels following glucose ingestion. Anthocyanin-rich MqBEs also reduced G6Pase and increased insulin sensitivity. Glucose uptake was upregulated in L6 skeletal muscle cells, without toxic effects [220].
Delphinidin 3-sambubioside-5-glucoside, a signature MqB biomarker, showed an equivalent capacity to metformin for normalizing blood glucose levels [326]. Lipid accumulation was inhibited by 4–11% by MqBE treatment in 3T3-L1 mice; however, lipogenesis was inhibited by 6–38% during adipocyte differentiation. The lipogenesis inhibitor protein, preadipocyte factor 1, was upregulated in MqB-treated 3T3-L1 mice. MqB supplementation also exerted an anti-inflammatory response by reducing ROS expression by 9.8 to 61.8%. The expression of COX-2 and production of PGE2 was also evaluated in the RAW 264.7 macrophages to understand the anti-inflammatory mechanism of MqB. MqB inhibited PGE2 expression and reduced COX-2 expression (by 16.2–62%), inhibiting LPS-induced iNOS/NO production and COX-2/PGE2 pathway activation in macrophages [218,326]. MqB delphinidin anthocyanins inhibited glucose uptake and transport from the rat duodenum by inhibiting SGLT-1. The inclusion of MqB-derived 35% anthocyanins and 25% delphinidin glycosides in a rice-chicken diet effectively reduced postprandial glucose levels. Purified delphinidin anthocyanin supplementation with a normal diet reduced fasting glucose and insulin levels [218,326]. MqB anthocyanins, in capsular form (3 x 150 mg per day), decreased oxidized LDL-C and 8-isoprostaglandin F2α, a urinary excretion oxidative stress marker [221,326]. Furthermore, MqB-derived delphinidin treatment effectively increased AMPK phosphorylation. Gene expression analysis showed that sodium palmitate exposure upregulated lipid accumulating genes such as SREBF1, CPT1-A, patatin-like phospholipase domain containing 2, and FASN, which were reduced by delphinidin treatment. Delphinidin supplementation limited weight gain in HFD-fed C57BL/6Nhsd mice, but not increased liver weight. Glucose homeostasis variations induced by HFD/HCD were also minimized by delphinidin treatment [222]. Hidalgo et al. [327] showed that delphinidin supplementation in rat jejunum tissues/cells reduced the short circuit current generated by glucose addition to an Ussing chamber. Delphinidin halted 3-O-methyl-glucose incorporation in the mouse intestine, with effects similar to the inhibition of electrogenic glucose transportation by SGLT-1 [328]. In response to delphinidin treatment and FFA1 activation, the Gaq/11 subunit was coupled with inositol trisphosphate, propionyl l-carnitine, and diacglycerol upregulation, which modulates intracellular Ca2+ from the endoplasmic reticulum. In previous studies, delphinidin treatment also caused intracellular Ca2+ release and prevented 3-O-methyl-glucose uptake by FFA1 activation. Therefore, delphinidin may represent a new ligand class that can reduce intestinal glucose uptake through FFA1 activation and increased cAMP expression [327].

MqB juice (MqBJ) consumption limited oxidation in human subjects (Table 2). The copper-triggered LDL-C oxidation time lag increased with MqBJ consumption because anthocyanins chelate copper. LDL-C oxidation time is proportional to the MqBJ anti-oxidative capacity. H2O2 treatment-induced increased oxidative stress was reduced by MqBJ treatment in human umbilical vein endothelial cells [329]. A pilot study showed that the daily MqBE consumption with folic acid and berberine effectively reduced TC, LDL-C, oxidized cholesterol glycemia, free radical levels, and increased serum antioxidant capacity. Furthermore, the insulinemia, microalbuminuria, HDL, CRP, and TG values increased. MqB treatment counteracted hyperlipidemia, hyperglycemia, and ROS production in metabolic syndrome patients. An MqB polyphenol-based nutraceutical reversed low-grade-inflammation, oxidative stress, and atherosclerogenesis in pre-diabetic patients [223]. MqBE and purified anthocyanin consumption showed positive outcomes for post-stroke stress and depression in diabetic mice. MqBESs and anthocyanins can mitigate anhedonia in humans. Anhedonic mice consumed less sucrose with increased water intake, which was mitigated by MqBE or purified anthocyanins in a dose-dependent-manner. Stroke and stress biomarkers such as TBARS, SOD, CAT, and GSH levels decreased following MqBE/anthocyanin treatment in stroke model mice [219].

16. Conclusions

This review aimed to collect and discuss scientific evidence regarding the positive role of berry consumption on the prevention of diabetes and its complications. Available human, animal, and in vitro studies were collected and comprehensively presented. This review demonstrated that berry product consumption represents a reliable and effective method for preventing and managing metabolic hyperglycemic and hyperlipidemic conditions. Variations in postprandial glucose and
Conflicts of Interest: The authors declare no conflicts of interest.

Abbreviations

1-Deoxynojirimycin (1-DNJ); Acai berry (AB); acetyl coenzyme A carboxylase (ACC); acyl-CoA oxidase (ACOX); adhesion molecules nuclear factor (lIβα); advanced glycation end-product (AGE); albumin retention (AR); alcohol-free fermented blackberry juice (AFBBJ); alkaline phosphatase (ALP); aminotransferase (ALT); AMP-activated protein kinase (AMPK); angiotensin I-converting enzyme (ACE-1); apolipoprotein A (Apo)A-I; aspartate aminotransferase (AST); ATP-binding cassette (ABC); ATP-binding cassette transporter (ABCA1) Bilberry juice (BB); BB/BB extracts, (BBEE); Bilberries, (BBs); Black currant (BCT); BIB extracts (BIBE); BIB juice (BIB); Blueberries (BBs); body weights (BW); brown adipose tissues (BAdT); capillary albumin filtration (CAF); carbohydrate response element-binding protein (ChREBP); cardiovascular disease (CVD); carnitine palmitoyl transferase-1 (CPT-1); Cranberries (CrBs); cranberries juice (CrB-JSB); CrB extracts (CrBE); CrB extracts (CrBEs); CrB juice (CrBJ); C-reactive protein (CRP); Diabetes mellitus (DM); diabetic retinopathy (DN); diacylglycerol acyltransferases 2 (DGAT2); fasting blood sugars (FBS); fatty acid synthase (FAS); Food and Agriculture Organization of the United Nations (FAO); forkhead box O1 (FOXO1); free fatty acid (FFA); gamma-glutamyltransferase (γ-GT); gastrointestinal-digested BB slurry (GIDBBB); GB polysaccharides (GBPS); glucagon-like peptide-1 (GLP-1); glucose tolerance test (GTT); glucose transporter (GLUT4); glucose transporter 2 (GLUT-2); glucose-6-phosphatase, (G6Pase); glucose-stimulated insulin secretion (GSIS); glutathione (GSH); glycerol-3-phosphate acyltransferase (GPA1); glycogen synthase (GYS1); glycogen synthase 2 (GYS2); Goji berry (GB); high-carbohydrate diets, (HCD); high-fat diets, (HFD); high-sensitivity CRP (hs-CRP); Human aortic endothelial cells, (HAECs); human serum albumin (HSA); inducible nitric oxide synthase (iNOS); insulin receptor substrate-1/2 (IRS-1/IRS-2); insulin resistance, (IR); intercellular adhesion molecule-
1 (ICAM-1); intestinal mucosal barrier dysfunction, (IMBD); lactate dehydrogenase (LDH); Lingonberry (LB); lipopolysaccharides (LPS); Low-calorie dried cranberry, (LCDC); low-density lipoprotein cholesterol (LDL-C); lyso-phosphatidylcholine (LPC); lyso-phosphatidylethanolamines, (LPE); malondialdehyde (MDA); manganese superoxide dismutase, (Mn-SOD); Maqui berries (MB); microbial-fermented blackberry metabolites (GMBB); microsomal TG transfer protein (MTP); mitochondrial transcription factor A (TFAM); monocyte chemo-attractant protein-1 (MCP-1); Mulberries (MBs); Na-glucose co-transporter 1 (SGLT-1); nitric oxide (NO); nitric oxides (NOs); nod-like receptor pyrin containing 3 (NLRP3); non-alcoholic fatty liver disease (NAFLD); paraoxonase-1 (PON-1); perilipin regulatory element (SCOA); soluble vascular cell adhesion molecule (VCAM); soluble vascular cell adhesion molecule-1 (sVCAM-1); sterol regulatory element-binding protein 1c (SREBP-1c); stereotactotin (STZ); Toll-like receptors, (TLR); total cholesterol (TC); total glyceroldehyde (TG); Trolox equivalent antioxidant capacity (TEAC); Type 1 diabetes mellitus, (T1DM); type 2 diabetes mellitus, (T2DM); unsweetened dried CrBs (USCB); white bread (WB); World Health Organization, (WHO).

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