Nutraceuticals as a New Approach in Management of Diabetic Nephropathy

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Abstract

Diabetic nephropathy (DN), a fatal diabetic complication, is a prime reason of end-stage renal disease (ESRD), which is pathologically identified by thickened tubular basal and glomerular membranes, amassed extracellular matrix (ECM), and developing meningeal hypertrophy. Various deformities in the signaling pathway can interact to give rise to these pathologic activities in DN. In spite of the accessibility of numerous approaches to prevent these metabolic changes, which include proper diet, exercising regularly, control of weight and control of drugs, epidemiological data are observing the growing tendency of the complication, indicating both the multifactorial nature of these disorders as well as the scarce conformance of patients to begin strategies. A number of Nutraceuticals utilized in clinical practice were shown to aim the pathophysiology of diabetes mellitus, metabolic disorder and their complications and to favorably modify numerous biochemical and clinical endpoints. These compounds comprise of antioxidants, vitamins like vitamin C, E, D; Omega 3 fatty acids, alpha lipoic acid (ALA), dietary fibers, flavonoids, phytoestrogens and minerals like chromium, magnesium. Various areas of concern prevail with regard of the use of nutraceuticals and dietary supplements in this setting, inclusive of standardization of products, potential side effects, definition of dosing regimen, interaction of drugs and need for evidence based interactions.

Keywords: Diabetic Nephropathy, Nutraceuticals, Pathogenesis, Complications, omega-3-fatty acid, vitamin D, dietary fibers, curcumin, resveratrol, quercetin, berberine, vitamin C.

Introduction

Diabetes is a non-communicable metabolic disorder that arises due to hormonal imbalance such as insulin, glycogen. Two types of diabetes Type 1 diabetes alias insulin dependent diabetes mellitus (IDDM), is caused by lack of insulin secretion by beta cells of the pancreas and Type 2 diabetes alias non-insulin dependent diabetes mellitus (NIDDM), is caused by decreased sensitivity of target tissues to insulin [1]. Diabetes is widely recognized as an emerging epidemic that has a cumulative impact on almost every country, age group, and economy across the world. According to the International Diabetes Federation, it was found that in 2015, approximately 415 million people were suffering from diabetes worldwide, and this number is expected to exceed 640 million by the year 2040. In 2020, according to the International Diabetes Federation (IDF), 463 million people have diabetes in the world and 88 million people in the Southeast Asia region. Of this 88 million people, 77 million belong to India. According to the 2019 National Diabetes and Diabetic Retinopathy Survey report released by the Ministry of Health and Family Welfare, the prevalence was found to be 11.8% in people over the
The prevalence of diabetes is 6.5% and prediabetes 5.7% among the adults below the age of 50 years, according to the DHS survey [2]. There are various complications that are associated with long-term damage and failure of various organ systems. Diabetes complications are common in patients with type 2 diabetes. The chronic complications of diabetes are broadly categorized into microvascular and macrovascular, with the former having much higher ascendency than the latter. Microvascular complications include neuropathy, nephropathy, and retinopathy, while macrovascular complications consist of cardiovascular disease, stroke, and peripheral artery disease (PAD) [3].

Diabetic Nephropathy (DN) alias Diabetic Kidney Disease (DKD) is a dreadful impediment of type 1 and type 2 diabetes mellitus (DM) associated with the kidneys [4]. DN is a medical condition depicted by glomerular hypertrophy, protein in urea, diminished glomerular filtration and kidney fibrosis which further bring about deprivation of kidney function [5]. DN is proclaimed to eventuate in 20%-50% of individuals living with DM and is the prevailing root of End Stage Kidney Disorder (ESKD) [6]. Indications of general DN is perpetual albumin in urea with retinopathy and no authentication of alternative kidney disorder [7]. The possibility of evolving DN differs between individuals and it relies not only on span of DM but is influenced by numerous other agents. Type 2 DM reports about 90% of diabetes throughout the world, so the large number of individuals who suffered from DN is because of type 2 DM [8]. Diabetic nephropathy can be rectified by correcting these parameters like Cardiovascular risk alleviation, maintaining glycaemic level, maintaining blood pressure, Blocking of Rennin Angiotensin Aldosterone System (RAAS), Changing the way of life [9]. Some of the allopathic drugs used in the treatment of diabetic nephropathy are Keresan, Benzoztamine, Robotisation, Telmisartan, Rapamycin, Fenofibrate, Liseran, and Barnidipine [5]. Allopathic drugs may cause long term side effects but still used widely. Nowadays, nutraceuticals have received high advantage due to potential nutritional, safety, and therapeutic effects. The nutraceuticals are used widely, also may be due to the fact that many of them are easily available, inexpensive and available in small doses. They usually do not cause repulsive side effects [10]. The purpose of this review is to summarize nutraceuticals as a current approach in the treatment of Diabetic Nephropathy. According to our hypothesis if the combination of two or more nutraceuticals can be more effective as compared to single nutraceutical. Further clinical studies must be done to clarify the effect of combination of nutraceuticals.

**Diabetic Nephropathy Pathogenesis**

There are several pathways that may cause Diabetic nephropathy (DN), such as hyperglycemia-induced renal hyper filtration and renal injury, AGEs-induced increased oxidative stress, activated PKC-induced increased production of cytokines, chemokine’s, and different inflammatory and apoptotic signals. Among all, oxidative stress has been suggested to play a major role in the propagation of DN. It stimulates several signaling pathways involved in DN, like AGEs, PKC cascade, JAK/STAT signaling, MAPK, m-TOR, and SMAD [41].

**Polyol Pathway** is a process that increases the ratio of NADH/NAD and may result in both oxidative stress and activation of protein kinase C. The Polyol pathway involves two enzymes. (I) Aldose Reductase (AR), reduces glucose to sorbitol by using NADPH as a co-factor, and (ii) sorbitol dehydrogenase (SDH) that converts sorbitol to fructose with the aid of its cofactor NAD+. Then, the excess sorbitol is oxidized to fructose [42]. This flux of glucose through the Polyol pathway would increase Advance Glycation End Products (AGE) formation that may result in oxidative stress [43].

**AGEs pathways (Advance Glycation End Products):** - AGEs accumulate at site of micro vascular injury in diabetes, including the kidney [44], the retina and within the vasculature [45]. AGE stimulate
activation and expression of IL-6 and TGFβ1 via NF-kB dependent pathways [45]. The main site for reabsorption of filtered AGEs is proximal tubule [46]. TGF-β1 expression is significantly linked to accumulation of AGEs in the kidney [47]. AGEs lead to the transcriptional up-regulation of TGF-β1, possibly via PKC or oxidative stress. In experimental diabetes, oxidative stress is increased in proportion to the accumulation of AGEs [48]. This pathway can also lead to enhanced formation of free radicals both directly through catalytic sites in their molecular structure [49]. The generation of superoxide may be enhanced due to mitochondrial dysfunction induced by AGEs and carbonyl intermediates [50]. AGE also contribute to the release of pro-inflammatory cytokine and growth factor expression and adhesion molecule such as VEGF and CTGF, TGF-β1, IGF-1, PDGF, TNF-α, IL-1β, and IL-6 [44,50].

Figure 1. Schema outlining various factors in the Pathogenesis of Diabetic Nephropathy

AGEs= advanced glycation end products; ROS= reactive oxygen species; RAA= Renin angiotensin - aldosterone system; DAG= diacylglycerol; PKC= protein kinase C; NO= nitrogen oxide; mTOR =mammalian target of rapamycin; VEGF= vascular endothelial growth factor;

Protein kinase C pathway: - Out of eleven isoforms of PKC, nine PKCs are activated by DAG, which is formed from excess glyceraldehydes-3-phosphate. Increased glucose concentration leads to increase the amount of DAG, which activates PKC. PKC activation further leads to changes in renal blood flow [51], by decreasing production of NO [52], meningeal expansion, albuminuria and increases GFR, increases pro-inflammatory gene expression and vascular permeability in many models of experimental diabetes [53]. Activation of PKC may be responsible for the increased expression of ECM molecules both directly and through TGF- β1 over expression.

Hexamine Pathway: - The Hexokinase is an essential enzyme for conversion of fructose-6-phosphate into glucosamine-6-phosphate. The Glutamine: Fructose-6-Phosphateamidotransferase (GFAT) is the rate-limiting enzyme of this pathway. Both high glucose and Angiotensin II activates the GFAT promoter in meningeal cells (MC) [54] and further that may enhance flux through the Hexokinase. Over activation of GFAT in MC leads to increase both TGF- β and fibronectin expression [55].
**Involvement of pro-inflammatory cytokines in diabetic nephropathy:**

**TNF-α:** Renal hyper-filtration and hypertrophy are some of the early changes that occur in the kidney, which seems to be due to inflammation particularly by TNF-α. For TNF-α to exert its actions, two mediators responsible i.e., receptors [56] TNFR1 and TNFR2. As TNF-α binds to these mediators and results in the formation of the TNFR associated death domain (TRADD) [57]. IL-6 acts by binding to the membrane-bound IL6R and gp130 signal-transducing chain. This results in the auto phosphorylation as a result of which both JAKs closer to each other to phosphorylate one another inducing an intracellular signal and further activates transcription factors, signal transducer and activator of transcription (STAT)-3. This activation leads to the growth and proliferation of meningeal cells [58].

Once released, IL-6 binds to the membrane-bound IL6R and gp130 signal-transducing chain. This result in the auto phosphorylation and change in the conformational structure of Janus kinase (JAK) which brings both JAKs closer to each other to phosphorylate one another inducing an intracellular signal. This further phosphorylates and activates transcription factors, signal transducer and activator of transcription (STAT)-3. Consequently, this leads to the growth and proliferation of meningeal cells and subsequently a similar outcome as IL-1 as described previously [59].

**Role of Oxidative Stress in Diabetic Nephropathy**

Hyperglycemia-induced oxidative stress has been suggested as the common mechanism causing the cell damage seen in diabetic complications [60]. Oxidative stress plays a major role in pathological changes of the kidney [61]. An imbalance between Reactive Oxygen Species (ROS) and intracellular antioxidants leads to Oxidative stress [62]. In Hyperglycemia overproduction of superoxide induced by mitochondrial electron transfer chain is the major molecular mechanisms for diabetes. Furthermore, increased NADPH oxidase activity leads to production of ROS in diabetic nephropathy [63]. Moreover, activation of PKC pathway leads to the production of ROS in diabetes which is attenuated by PKC inhibitors.

**Genetic mechanisms leading to diabetic nephropathy**

**Angiotensin-converting enzyme (ACE):** The dysfunctional ACE gene can cause toxicity and fibrosis of blood vessels by increasing aldosterone level [65]. It was found that by the activation of the Smad2-dependent TGFb1 pathway the aldosterone aids in the production of the extracellular matrix protein, fibronectin by glomerular meningeal [66].

**FRMD3:** Another gene associated with DN is the FRMD3. An SNP, rs1888747 located on chromosome 9 in the promoter region of FRMD3 is strongly associated with DN [67] Martini and colleagues (2013) suggested that FRMD3 operates in conjunction with bone morphogenetic protein (BMP) signaling pathway [68]. BMP is part of the TGFb1 super-family plays a part in kidney development, chemotaxis, and cell differentiation and regulates apoptosis of various adult cell types such as hematopoietic, epithelial, neuronal and mesenchymal cells [69]. The rs1888747 SNP in the promoter region of FRMD3 is considered to suppress the activity of FRMD3, which prevents the activation of BMP and the depletion of BMP-mediated renal protection in patients with diabetes [70]. FRMD3/BMP inhibition can result in albuminuria and increase in fractional meningeal area. For example, BMP7 agonists and antagonists like kielin/chordin-like protein and gremlin respectively have been reported in patients with DN and suppressed the expression of BMP7 and its agonists have shown to increase profibrotic activity in DN [41, 42].
Nutritional Interventions Targeting the Pathogenesis of Diabetes Mellitus: -

Omega-3 fatty acids:

Omega-3 polyunsaturated fatty acids (n-3 PUFAs), including the essential fatty acids α-linoleic acid (ALA) and the long chain fatty acids eicosapentaenoic acid (EPA) and docosahexaenoic acid. It is an (DHA), obtained from plants and marine life. It gains great interest in the health benefits among the many chronic diseases in which n-3 PUFA intake can be improved, there is a large literature that discusses the therapeutic potential associated with type 2 diabetes [73]. The mechanism by which omega-3 fatty acids reduce proteinuria is not yet clear. There is evidence that omega-3 fatty acids may affect renal hemodynamic [74]. The effect of omega-3 fatty acids on increasing proteinuria may exceed hemodynamic parameters. One hypothesis is that omega-3 fatty acids may reduce urinary protein excretion via anti-inflammatory effects and oxidative stress [75]. O3FA also may reduce proteinuria in patients with chronic glomerular disease and delayed immunoglobulin A (IgA) nephropathy [76]. Because of its anti-inflammatory action, O3FA is thought to prevent kidney damage. An anonymous model study has shown that O3FA formulations reduce renal inflammation and fibrosis [77]. Experimental Diabetic Nephropathy Garman et al. studied the effect of omega-3 polyunsaturated fatty acids administered to male streptozotocin-induced diabetic Sprague-Dawley rats for 30 weeks. Mice that consumed polyunsaturated fatty acids high in nonomega-3 had significantly less albuminuria, high blood pressure, glomerulosclerosis, and interstitial tubular fibrosis. In addition, the increase in type IV and this collagen, IL-6, MCP-1, TGF and the macrophage marker CD68 was reduced. The researchers concluded that omega-3 polyunsaturated fatty acids significantly reduce kidney damage caused by diabetes [78].

Berberine:

Berberine (BBR; [C20H18NO4] +).an isoquinoline alkaloid isolated from the rhizome of Coptidis, cortex philodendra and Berberis vulgaris, has long been used widely as an Eastern medicine for the treatment of gastroenteritis and secretarydiarrhea [79-82]. Recent studies have illustrate that berberine has various pharmacological activities, including lowering blood glucose, regulating blood lipids and reducing inflammation in addition to its antioxidant activity. These findings suggest that berberine has applications as a therapeutic drug for diabetic nephropathy, and has significant research value [83]. In addition, BBR is well characterized as having a Reno protective effect on the development of DN [84, 85]. However, the possible mechanisms have not been fully established. The purpose of this article is to investigate the Reno protective mechanisms of berberine in diabetic nephropathy and highlight the importance of berberine as a potential therapeutic reagent for diabetic nephropathy treatment [83]. Inflammation, fibrosis, and lipid abnormalities are the main causes of the pathogenesis of diabetic kidney damage in type 2 diabetes.[86] Berberine (BBR) has been reported to have beneficial effects in diabetic nephropathy, but its mechanism of action is unclear. This study aims to determine the therapeutic mechanism of LBW in a rat model of type 2 diabetic nephropathy induced by a high-fat streptozotocin regimen and low-dose injection. Diabetic mice were given tubes with or without BBR for 20 weeks and examined by serological, 24-hour proteinuria, histology, immunohistochemistry, and molecular tests [86]. The results showed that LBW treatment significantly reduced serum sugar and lipids, inhibited urinary tract albumin excretion, and attenuated histological kidney damage in diabetic rats. Berberine treatment also suppresses renal inflammation associated with inactivation of a key factor that increases kappa through an activated B-cell signaling light chain. As a result, the up regulation of pro-inflammatory cytokines (interleukin-1β, tumor necrosis factor-α) and chemokine’s (protein of chemotactic monocytes-1) is blocked. In addition, BBR treatment inactivates growth
factor-β / Smad3 transforming signal modification and suppresses renal fibrosis, including the expression of fibronectin, collagen I, and collagen IV. This study shows that BBR is a diabetes remedy. Relief of type 2 nephropathy consists in suppressing the kappa-nuclear potentiator of the activated inflammatory light chain of B cells and altering the signaling pathway of growth factor-β / Smad3 [85].
# Table no. 1 List of Nutraceuticals used in Diabetic Nephropathy

<table>
<thead>
<tr>
<th>Nutraceuticals</th>
<th>Source</th>
<th>Mechanism of action in DN</th>
<th>Other activities</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Omega-3 fatty acids</td>
<td>obtained from plants and marine life</td>
<td>may reduce urinary protein excretion through anti-inflammatory effects and oxidative stress</td>
<td>Anti-inflammatory, anti – thrombotic, decrease TGs, reduce blood clotting time</td>
<td>Clark WF et al</td>
</tr>
<tr>
<td>Berberine</td>
<td>isoquinoline alkaloid isolated from the rhizome of Berberis vulgaris</td>
<td>Reno protective by reducing fibrosis, Inflammation, and lipid abnormalities</td>
<td>lowering blood glucose, regulating blood lipids and reducing inflammation</td>
<td>Han J et al</td>
</tr>
<tr>
<td>Dietary fibers</td>
<td>obtained from plant products of polysaccharides, oligosaccharides, lignin</td>
<td>Modulation of gut micro biota, enrichment of SCFA-producing bacteria</td>
<td>increasing body weight, lowering blood cholesterol levels and, coronary artery disease, dyslipidemia and hypertension</td>
<td>JULIE M J et al</td>
</tr>
<tr>
<td>Quercetin</td>
<td>Cabbage, onions, fruits, apple, red wine, broccoli, cherries, tea and red wine etc.</td>
<td>powerful antioxidant properties potent ROS scavenger and may reduce the risk of cardiovascular disease</td>
<td>prevents various diseases such as osteoporosis, some forms of cancer, tumors, lungs,</td>
<td>Chen T.C. et al</td>
</tr>
<tr>
<td>Alpha lipoic acid [ALA]</td>
<td>spinach, then broccoli, tomatoes, garden peas and rice bran</td>
<td>Powerful ROS scavenger, renew endogenous antioxidants</td>
<td>the treatment of Alzheimer’s disease, cancer, diabetic complications, multiple sclerosis, diabetes, obesity</td>
<td>Oza M.J. et al</td>
</tr>
<tr>
<td>Vitamin D</td>
<td>flesh of fatty fish and fish liver oils egg yolks, cheese, and beef liver</td>
<td>encouraging insulin production, restrict renin production, and stimulating macro-phage cathelicidin development</td>
<td>Renal protective by proteinuria, antifibrosis, anti-inflammatory, and Inhibiting podocyte devastation</td>
<td>Holick, M. F. et al</td>
</tr>
<tr>
<td>Flavonoids</td>
<td>fruits, vegetables, tea, cocoa and wine, onion, tea</td>
<td>Anti- oxidant</td>
<td>anti-bacterial, anti-hypertensive, anti-diabetic, anti-inflammatory, anti-Parkinson, anti-ulcer, spasmyolytic, anti-depressant</td>
<td>Havsteen B et al</td>
</tr>
<tr>
<td>Resveratrol (3,4',5-trihydroxystilbene)</td>
<td>mainly found in grapevines and berries</td>
<td>reducing oxidative stress and advanced glycation end-product (AGE) production</td>
<td>stimulating autophagy, inhibiting endoplasmic reticulum (ER) stress and inflammation, ameliorating lipotoxicity, activating the AMP kinase (AMPK) pathway, and modulating angiogenesis</td>
<td>Evans JL et al</td>
</tr>
<tr>
<td><strong>Curcumin</strong></td>
<td>extracted from the rhizome <em>Curcuma longa</em></td>
<td>activates various transcription factors, such as NF-E2-related factor (Nrf2), per-oxisome proliferator-activated receptor-γ (PPAR-γ), CCAAT/enhancer binding protein (C/EBP) homologous protein (CHOP)</td>
<td>treat diabetic nephropathy, microangiopathy and retinopathy</td>
<td>Khajehdehi et al 221</td>
</tr>
<tr>
<td><strong>Vitamin C</strong></td>
<td>Citrus fruits and vegetables like lemon, orange etc.</td>
<td>chain-breaking antioxidant, scavenging ROS directly, and preventing the propagation of chain reactions</td>
<td>decreasing lipid peroxidation and augmented the activities of antioxidant enzymes, SOD, and GPx and reduced albuminuria</td>
<td>Riccioni G et al</td>
</tr>
</tbody>
</table>

TGs= Triglycerides; ROS=reactive oxygen species; SCFA= short chain fatty acid; ER= endoplasmic reticulum; AMP= adenosine monophosphate; Nrf2= nuclear factor erythroid-2-related factor; SOD= superoxide dismutases; GPx= glutathione peroxidase

**Dietary fibers**

The term "dietary fiber" refers to a wide variety of natural food sources, refined grains, and commercial nutritional supplements. Dietary fiber is "an edible part of a plant or similar carbohydrate that fights digestion and digestion in the small intestine through complete or partial fermentation in the human large intestine. It is obtained from plant products of polysaccharides, oligosaccharides, lignin and other substances [86, 87, and 88]. Pharmacological use of dietary fibers: increasing body weight, lowering blood cholesterol levels and / or lowering blood sugar levels, coronary artery disease, dyslipidemia and hypertension [86]. Several studies have shown that the US population and diabetics do not respond to adequate daily fiber intake from their diet [88]. Mechanism of action dietary fibers in case of diabetes nephropathy. **Soluble fiber:** has been linked to decreased postprandial glucose levels and increased insulin sensitivity in diabetics and healthy people. This effect is usually associated with the viscous and / or gelling properties of soluble fiber [87, 88]. Insoluble fiber: Does not significantly affect post-meal blood sugar. While, consuming soluble fiber does not reduce the risk of type 2 diabetes [60]. In contrast, consumption of insoluble dietary fiber was most closely associated with a reduced risk of diabetes. To study the effect of fiber on the development of experimental diabetic nephropathy, previously experimental in which streptozotocin to induce diabetes in C57BL / 6 wild and knockout mice lacking the gene encoding the G-protein coupled receptor GPR43 or GPR109A. Diabetic mice were randomly assigned a diet high in fiber, normal, less in fiber, or high in SCFA in their drinking water. They found proton magnetic resonance imaging techniques to determine the metabolic profile and 16S ribosomal RNA sequencing to determine the entire gut micro biota [89]. Diabetic mice fed a high-fiber diet were significantly less likely to develop diabetic nephropathy. Mild proteinuria, glomerular hypertrophy, podocyte injury, and interstitial fibrosis were compared with control diabetic patients who received a normal or fiber-free diet. Dietary fiber promotes the proliferation of SCFA-producing bacteria from the genus Prevotella and Bifid bacterium, positively altering the microbial ecology of the intestine and improving dysbiosis by increasing the levels of SCFA in the faeces and the system. Reduction of fiber
expression of genes encoding inflammatory cytokines, chemokine’s, and proteins that induce fibrosis in diabetic kidney. Diabetic mice treated with SCFA were protected from nephropathy, but not in the absence of GPR43 or GPR109A. SCFA in vitro regulates inflammation of renal tubule cells and paw cells under hyperglycemic condition [89].

**Quercetin**

Quercetin is a flavonoid compound and it is belonging to a group of plant derived heterocyclic polyphenols [90]. It is obtained from in large amount of Cabbage, onions, fruits, apple, red wine, broccoli, cherries, tea and red wine etc. [91, 92]. Recent research shows that biochemical and pharmacological studies of Quercetin have shown that it is a potent ROS scavenger and may reduce the risk of cardiovascular disease and kidney disease [90, 93-95]. Quercetin also prevents various diseases such as osteoporosis, some forms of cancer, tumors, lungs, and diabetes mellitus. The antioxidant properties of quercetin play an important role in the prevention and treatment of these diseases [96]. Mechanism of action quercetin in case of Diabetes nephropathy (DN): Accumulating evidence suggests that increased oxidative stress may play an important role in the pathogenesis of DN [97, 98], but antioxidant therapy in diabetic patients has shown conflicting results during DN treatment [99]. Currently, only blockers of the renin-angiotensin system are chronically used. However, this mechanism has limitations [99,100], which justify the search for safer and more effective candidates for antioxidants. Flavonoids and their relative product i.e., quercetin have powerful antioxidant properties and are plant derived phenolic compounds found throughout the plant kingdom. Hence, Flavonoids are a therapeutic option [101,102]. Collected 8-week-old homozygous mice C57BL / 6J. Animals were placed at room temperature (22° C) in a controlled humidity environment with a 12 hour / 12-hour dark light cycle in a laboratory translational physiology laboratory apparatus. Diabetic mice by intraperitoneal injection of STZ, diluted in citrate buffer (10 mm, pH 4.5) at a dose of 100 mg / kg / day for 3 days. Controls (non-diabetic, ND, n = 10) were given an equivalent volume of citrate buffered vehicle. Blood glucose was measured after food shortage for 6 hours after 1 week of STZ injection, and animals with glucose levels of 250 mg / dL or more for a minimum of 2 days were considered diabetic. Six weeks after STZ injection, diabetic mice were randomized to receive no treatment (soybean oil carrier, DV, n = 10) or oral quercetin (DQ, n = 10) at 10 mg / kg per day for 4 weeks. This dosage is based on previous studies on hypertension and diabetics animals [103,104]. This study suggests that chronic oral treatment with low doses of quercetin has an antidiabetic effect and attenuates the development of nephropathy in STZ-induced DN mice. These results are supported by reductions in plasma glucose, creatinine, triglycerides, proteinuria, and decreased interstitial matrix proliferation that accompanies. O2 production and apoptosis in kidney cells. In mice injected with STZ, β-cell damage and a decrease in the ability to secrete insulin were found [105,106]. These results support the role of oxidative stress in the development of diabetic nephropathy and suggest a possible antioxidant mechanism responsible for the nephroprotective effect of quercetin.

**Alpha lipoic acid [ALA]**

Alpha lipoic acid (ALA) is an anti-oxidant which occurs naturally in foods covalently bound to lysine in proteins [107]. Alpha lipoic acid is of two types: one is the R- alpha lipoic acid and the other is S-alpha lipoic acid. The R -isomer is present in nature and S-isomer is not present in nature [108]. The most copious plant source of ALA is spinach, then broccoli, tomatoes, garden peas and rice bran [109]. Extreme concentrations of ALA are observed in animal tissues with immense metabolic action namely heart, liver and kidney. Contrary to ALA in foods, ALA in supplements is unbound to protein.
Furthermore, the proportion of ALA available in dietary supplements (200–600 mg) is expected to be 1000 times greater as compared to the proportion that could be acquired from the diet [107]. Four antioxidant characteristics of ALA have been analyzed: it’s metal chelating activity; it’s potential to scavenge reactive oxygen species (ROS); it’s capacity to renew endogenous antioxidants and its property to restore oxidative destruction [110]. ALA is clinically used in the treatment of Alzheimer’s disease, cancer, diabetic complications, multiple sclerosis, diabetes, obesity and also it is used as an antioxidant [111]. Oxidative stress seems to depict a chief role as a rapid development factor in diabetic nephropathy (DN) and later advancement [112]. ALA has powerful ROS-scavenging capacity. It has the exceptional attribute of being a ROS scavenger in its oxidized condition, quenching several radicals. ALA and dihydrolipoic acid (DHLA) together work as a redox couple (an electron donating molecule and its oxidized form), and have additional antioxidant characteristics comprising chelation of transition metals and the renewal of other antioxidants glutathione, vitamin C, and vitamin E, for instance [7,8]. ALA serve as a helper molecule for Various key mitochondrial multienzyme complexes. Improve the consumption of glucose by the cells, and modify the action of numerous signaling molecules and transcription elements. ALA and its derivative, DHLA, have a Straight antioxidant property owing to the neutralization of reactive oxygen species that are devastating to DNA, proteins and lipids of cells [115]. Oxidative stress gives rise to endothelial cell (EC) damage and vascular dysfunction [116].

In this study, 49 patients (34 with type 1 diabetes and 15 with type 2 diabetes) do not undergo antioxidant therapy and we’re said to be standard. The remaining 35 patients (20 with type 1 diabetes and 15 with type 2 diabetes) Undergo ALA therapy (600 mg/day for 18 months). The development of EC Destruction with regards to the evaluation of plasma thrombomodulin was remarkably increased in the standard group and reduced in the ALA therapy group after 18 months of research. However, the advancement of DN, as estimated by urinary albumin level, was considerably elevated in standard, but remains unaffected in the treated group. [117]

**Vitamin D**

Vitamin D (VD) is very uncommon in unfortified foods. VD is classified in two categories: vitamin d2 and vitamin d3. VD in different proportions is present in the flesh of fatty fish and oils of fish as well as cod and tuna liver oil [118,119]. Various foods are vitalized with Involving milk, few cereals and few bread items. There is preparatory authentication to indicate that meats from poultry, pork and beef comprise little proportion of VD that possibly eventuates from the VD that was vitalized in the animal feed [120]. Milk is also regarded as to be the food source of VD [121]. The vital source of VD for nearly all human beings is natural subjection to sunlight. It is determined that 80-90% of the body's demand for VD is fulfilled from this source [119]. Vitamin D receptor (VDR) emerges in various tissues and cells in the body [122,123]. 1,25(OH)2D has a broad extent of pharmacological Properties, namely hindrance of cellular proliferation and causing terminal differentiation, hindrance of angiogenesis, encouraging insulin production, restrict renin production, and stimulating macrophage cathelicidin development[122-125]. The deprivation of VD is the root cause of numerous chronic disorders like osteoporosis, fractures, diabetes, autoimmune disease, hypertension, periodontal disease, multiple sclerosis, cognitive impairments, Parkinson’s disease, osteoarthritis, bacterial vaginitis. So, to prohibit all these disorders, the level of vitamin d should be accordingly maintained [126]. VD, a type of steroid having operative form 1, 25(OH)2D3, has been well known for the Significant functions in the modulation of serum calcium and phosphorus concentrations. It provides salient features by binding with its receptor VDR. VDR, a transcription agent present at chromosome 12 comprising 9 axons, is distinct nonsteroid nuclear hormone receptor super family, which gets involved in transcriptional management of genes in tissue- and cell-specific ways.
Increasing documentation have indicated that VD/VDR signaling pathway procure a range of renal-protective outcomes in DN patients, including proteinuria, antifibrosis, anti-inflammatory, and Inhibiting podocyte devastation [127]. Animal studies have revealed that serum 25(OH) D (calcifediol) and 1, 25(OH) 2D3 (calcitriol) levels were considerably diminished in the DN rats and VD therapy of the rat model efficaciously prevented hyperglycemia-induced meningeal cell proliferation [128]. In one more animal study, diabetic VDR-/- mice exhibited more acute kidney injury and were expected to have severe proteinuria and glomerulosclerosis in contrast to the wild-type mice [129].

**Flavonoids**

In nature, Flavonoids compounds are products obtained from plants and they are detected in various parts of the plant. Flavonoids are utilized by vegetables for their development and Protection against plaque [130]. Plant based food and beverages comprise of plentiful of flavonoids namely fruits, vegetables, tea, cocoa and wine; therefore, they are named as dietary flavonoids. Flavonoids have various subdivisions, which contain chalcones, flavones, flavanols and isoflavones. These subdivisions have distinctive principal sources; onions and tea are chief dietary sources of flavanols and flavones, for instance [131]. Beside various biological properties in plants (resistance to herbivores, UV radiation and pathogens), flavonoids carry out countless pharmacological properties viz; anti-Parkinson, anti-ulcer, spasmolytic, anti-depressant, anti-bacterial, anti-hypertensive, anti-diabetic, anti-inflammatory and anti-cancer [132]. New insights are proposed for the probable utilization of flavonoid derivatives as beneficial medicine to control specific cancers [133]. Flavonoids also provide auspicious benefits obesity- and inflammation-as well as COVID-19 [133]. Scientific has strongly claimed that regular consumption of dietary flavonoids in potent amounts minimize the possibility of oxidative stress and chronic inflammation-mediated pathogenesis of human diseases such as CV disorder, specific cancers, neurological diseases and nephropathy [134]. Diabetic nephropathy (DN) is one of the most severe complications in DM, and it could be the reason of end-stage kidney disorder among sufferers which underwent chronic haemodialysis treatment [135]. In combination with a good hypoglycemic activity, flavonoids can be used to inhibit the occurrence and advancement of nephropathy [136]. The flavonoid mixture silymarin was estimated for its effectiveness on oxidative stress and morphology of kidney tissue in alloxan-induced diabetic rats. Tissue Devastation due to alloxan was inhibited by silymarin therapy and the activity and gene expression of three antioxidant enzymes were reinstated after the therapy with silymarin [136]. In further investigation, the preventive effect of silybin against high glucose-induced podocyte injury was discovered manifesting its Use against DN [137]. Countless investigations have illustrated antioxidant effects of various flavonoids on kidney. Depleted oxidative stress load and lessened renal devastation in a type 2 diabetic rat model by increasing the activity of the antioxidant enzymes Superoxide dismutase (SOD) and catalase (CAT), accompanied by introduction of the protective molecule Sirtuin-1 [138]. *Diosmetin*, a flavonoid extracted from the leaves of oleaeuropaea with powerful antioxidant property, reinstated the lower levels of SOD and Nitric Oxide (NO) observed in the kidney of diabetic rats and diminished the levels of malondialdehyde (MDA), a lipid peroxidation marker [139].

**Curcumin**

Turmeric or Curcuma longa is a natural product, with a wide variety of therapeutic effects on several diseases such as neurodegenerative, hepatic and renal damage, cancer, and diabetes have been mainly attributed to its curcuminoid content. It is a bright yellow colored component extracted from the
rhizome *Curcuma longa*, commonly known as turmeric [140]. The current basic evidences about the potential of curcumin/curcuminoids for the treatment of diabetes mellitus, mainly by its hypoglycemic, antioxidant, and anti-inflammatory properties [141]. Modern scientific studies shows that Curcumin is a highly pleiotropic molecule that interacts with numerous molecular targets, which it can up- or down regulate, Curcumin activates various transcription factors, such as NF-E2-related factor (Nrf2), peroxisome proliferator-activated receptor-g (PPAR-g), CCAAT enhancer binding protein (C/EBP) homologous protein (CHOP) and activating transcription factor 3 and it also been shown to down regulate various transcription factors, including NF-kB, protein kinases, chemokine’s and inflammatory biomarkers [142]. A study showed that curcumin increases the half-life, mean residence time, and the apparent volume of distribution at steady state of glibenclamide in rats that may be due to decreased metabolism of the drug mediated by the inhibition of intestinal and hepatic CYP3A4 [143]. Despite the potential tremendous benefits, the clinical trials of curcumin are only available in using curcumin to treat diabetic nephropathy, microangiopathy and retinopathy so far. Clinical trials with curcumin in patients with overt type 2 DN (Diabetic nephropathy), a curcumin supplementation at a dose of 66.3 mg per day for 2 months attenuated proteinuria, TGF-b and IL-8 [144]. Curcumin supplementation at the same dose has also been proved to decrease proteinuria, hematuria and systolic blood pressure in patients with relapsing or refractory lupus nephritis, which suggests that it can be used as an adjuvant safe therapy for such patients [145]. Further, there are several approaches to overcome limited solubility and poor bioavailability, these include encapsulation in various nanoparticles, such as methionine–dehydrophenylalanine nanoparticles, N-trimethyl chitosan chloride-coated liposomes, chitosan micro particles and in Nano suspension emulsions, and sustained released tablets [146-148].

**Vitamin C (ascorbic acid)**

Vitamin C is also known as ascorbic acid, and act as a cofactor in multiple enzymatic reactions including collagen synthesis.8 Citrus fruits and vegetables are the major source for Vitamin C such as orange sand orange juice, strawberries, blackcurrants, peppers, , broccoli, Brussels sprouts, potatoes. For adults the recommended dietary intake of vitamin C is 45 mg per day. A wide variety of disorders like diabetes, atherosclerosis, common cold, cataracts, glaucoma, macular degeneration, stroke, heart diseases, and cancer and so on, can be treated and prevented by Vitamin C. Vit. C is a chain-breaking antioxidant, scavenging ROS directly, and preventing the propagation of chain reactions that would otherwise show reduction in protein Glycation [149]. The diabetic condition possibly causes a variety of tissue damage in patients with diabetes and also production of oxidative stress. And further, the increased oxidative stress in the diabetic kidney may induce apoptosis, which may contribute to the development of diabetic nephropathy [150,151]. Vitamin C plays a crucial role in the antioxidant defense system and in the apoptosis. It was demonstrated that vitamin C exclusion from tubular epithelial cells, through the competition of glucose and dehydroascorbate for a common transport mechanism, will deprive the cells of antioxidant ability and could lead to ROS accumulation in diabetes [152]. It was demonstrated that in diabetic rats the administration of vitamin C can protect podocyte by increasing antioxidative capacity and ameliorating the renal OS [153,154]. In addition, it also acts by decreasing lipid peroxidation and augmented the activities of antioxidant enzymes, SOD, CAT, and GPx and reduced albuminuria, and GBM thickness in the kidneys of diabetic rats [155]. It was found that vitamin C protects renal lesions in DN by inhibiting expression of type IV collagen [156]. It has been shown that via Nrf2 activation vitamin C increases HO-1 protein expression in a concentration- and time-dependent manner. In animals, it brings down the diabetes-induced sorbitol accumulation and lipid peroxides in erythrocytes [149]. When vitamin C is combined with metformin a significant reduction in HbA1c levels in diabetic patients observed as compared to metformin alone.
Another study showed that administration of vitamin C with antidiabetic drugs enhanced antioxidant defenses and reduced oxidative damage [158].

**Resveratrol**

Resveratrol (3, 4’, 5-trihydroxystilbene) is a nutraceutical that is mainly found in grapevines and berries and available in various pharmaceutical dosages. It is a phytoalexin, firstly isolated from *Veratrum grandiflorum*. Resveratrol has nephroprotective effects through various mechanisms including reducing oxidative stress and advanced Glycation end-product (AGE) production, stimulating autophagy, inhibiting endoplasmic reticulum (ER) stress and inflammation, ameliorating lipotoxicity, activating the AMP kinase (AMPK) pathway, and modulating angiogenesis that eventually help in maintaining optimum kidney function during DN [159,160]. Moreover, the use of resveratrol as an adjuvant to conventional antidiabetic therapies could be an effective approach to manage DN. Previous reports suggest that elevated glucose flux can increase reactive oxygen species (ROS) production by the mitochondrial electron transport chain [161]. The accumulation of ROS can overwhelm the cellular antioxidant defense system and thus induce renal damage via different mechanisms, including activation of the polyol pathway and Hexosamine biosynthesis pathway, AGEs production, and activation of protein kinase C (PKC) [159,160]. Therefore, natural products like resveratrol with potential antioxidant capacity can attenuate these pathways and the associated complications for the control of DN. It has been founded that ER stress is one of the major contributing factors in the onset and progression of various pathological conditions including DN [162]. In addition, Resveratrol also acts by inhibition of renal epidermal growth factor receptors (EGFRs) and reduce ER stress in DN [163]. Pre-clinical studies shows that Resveratrol treatment [5 and 10 mg/(kg · d)] in a diabetic rat model (streptozotocin-induced DM), significantly attenuated pre-DN symptoms including reduced clearance of creatinine and urea, proteinuria, and oxidative stress (elevated lipid peroxidation and decreased antioxidant enzyme activity) via its antioxidant activity and resveratrol treatment [5 mg/(kg · d)] in type 1 diabetes rats significantly improve renal hypertrophy and structural changes, including tubular atrophy, meningeal expansion, diffused glomerulonephritis, and renal fibrosis. Preliminary human studies confirm the effectiveness of Resveratrol as given: - 613 type 2 diabetes confirm the effectiveness of this compound, 614 particular improves glycaemic control and 615 decreases insulin resistance [134,135,136].

**Conclusion:** Among many disease or disorders of carbohydrate, fat and protein metabolism, diabetes is most serious disorder effecting large population of the world. It is allied with decreased insulin production or resistance towards its action. The routine medical treatments used for DM are notenough therapeutic effective and have many undesirable side effects. Moreover, the global increased dispersal of DM makes researchers try to find out potential complementary or alternative treatments. Plants can be used to treat diabetes patients, both insulin dependent & non-insulin dependent diabetes. Nutraceuticals are the food supplements having nutritional value. All the nutrients discussed in this review have shown significant clinical & pharmacological activity. The potency of herbal drugs is prominent & they have very less side effects than the synthetic anti-diabetic drugs. There is increasing demand by patients to use the natural products with anti-diabetic activity. This review summarize the role of nutraceuticals in diabetic nephropathy (DN). The nutraceuticals including omega-3 fatty acid, vitamin D, dietary fibers, curcumin, resveratrol, quercetin, berberine, vitamin C have beneficial effects on pathophysiology of DN. Most of the nutraceuticals used in clinical practicemodulate one or more pathogenic mechanismsunderlying the progress of diabetes mellitus, metabolic syndrome, and their complications.
References


