Review Article

Bioactive Constituents and Toxicological Evaluation of Selected Antidiabetic Medicinal Plants of Saudi Arabia

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The purpose of this review is to summarize the available antidiabetic medicinal plants in the Kingdom of Saudi Arabia with its phytoconstituents and toxicological findings supporting by the latest literature. Required data about medicinal plants having antidiabetic activities and growing in the Kingdom of Saudi Arabia were searched/collected from the online databases including Wiley, Google, PubMed, Google Scholar, ScienceDirect, and Scopus. Keywords used in search are in vivo antidiabetic activities, flora of Saudi Arabia, active ingredients, toxicological evaluations, and medicinal plants. A total of 50 plant species belonging to 27 families were found in the flora of Saudi Arabia. Dominant family was found Lamiaceae with 5 species (highest) followed by Moraceae with 4 species. β-Amyrin, β-sitosterol, stigmasterol, oleanolic acid, ursolic acid, rutin, chlorogenic acid, quercetin, and kaempferol are the very common bioactive constituents of these selected plant species. This paper has presented a list of antidiabetic plants used in the treatment of diabetes mellitus. Bioactive antidiabetic phytoconstituents which showed that these plants have hypoglycemic effects and highly recommended for further pharmacological purposes and to isolate/identify anti-diabetes mellitus (anti-DM) active agents also need to investigate the side effects of active ingredients.

1. Introduction

Medicinal plants are used for the treatment of different infections [1, 2]. These plants contributed as a source of inspiration for novel therapeutic compounds [3]. The medicinal value of plants is due to the presence of a wide variety of secondary metabolites including alkaloids, glycosides, tannins, volatile oil, and terpenoids [4, 5]. Medicinal plants and their extracts represent a rich source of crude medications that possess therapeutic properties. Indeed, the World Health Organization reports that various plant fractions and their dynamic constituents are utilized as traditional medicines by 80% of the world population [6]. Plants are the primary source for different pharmaceutical, perfumery, flavor, and cosmetics industries; the use of modern drugs dramatically resulted into resistant microorganisms toward different modern drugs; the researchers are now in search for alternate source of treatment of various disorders [7, 8]. For this purpose, the medicinal herbs are the best alternate to various drugs. Most of natural products possess interesting biological activities and medicinal potential. Various herbs, fruits, and grains have been found to have different important biological activities such as antioxidant, [9] antitumor, antimitogenic, antidiabetes, antianalgesic, [10] antidegeneration, inflammation inhibitory effect, [9] antitumor, [11] anticancer, [12] antimicrobial, antileishmanial, and antimalarial properties [13, 14]. The consumption of natural antioxidants will reduce risk of many diseases including cancer, cardiovascular disease, diabetes, and other diseases allied with aging [15]. For natural antioxidants, a larger number of medicinal herbs have been evaluated by applying laboratories’ developed procedures. Plants derived substances, collectively called phytonutrients or phytochemicals, been recognized as good source of natural antioxidants [16, 17].

The Kingdom of Saudi Arabia is a huge arid land with an area of about 2,250,000 km² covering the major part of the Arabian Peninsula, characterized by different ecosystems...
and diversity of plant species. The climate in Saudi Arabia differs greatly between the coast and the interior. High humidity coupled with more moderate temperatures is prevalent along the coast, whereas aridity and extreme temperatures characterize the interior. The flora of Saudi Arabia is one of the richest biodiversities in the Arabian Peninsula and comprises very important genetic resources of crops and medicinal plants. Saudi Arabia contains 97 trees, 564 shrubs, and about 1620 herbs, which cover, respectively, 4.25%, 24.73%, and 71.02% of higher plant diversity of the country [18].

Diabetes mellitus is one of the most prevalent diseases in endocrine gland system with an increasing incidence in human community [19]. Type I diabetes is caused by insulin secretion deficit, while type II diabetes is accompanied with progressive rate of insulin resistance in liver and peripheral tissues, reducing β-cell mass, and deficient insulin secretion [20, 21]. This disease brings about acute metabolic side effects including ketoacidosis, hyperosmolar coma accompanied with chronic disorders, and long term, adverse side effects such as retinopathy, renal failure, neuropathy, skin complications, as well as increasing cardiovascular complication risks [22, 23]. Also, common symptoms of diabetes are frequent urine, thirsty, and overeating [24]. Diabetes inflicts 100 million people yearly and is recognized as the seventh cause of death in the world [25]. It has been estimated that the number of diabetic people will increase from 150 million individuals in 2003 to 300 million by 2025 [26].

The essential and effective drugs for diabetes mellitus are insulin injection and hypoglycemic agents, but these compounds possess several adverse effects and have no effects on diabetes complications in long term. Therefore, it is important to find effective compounds with lower side effects in treating diabetes [27].

Medicinal plants are good sources as alternative or complementary treatments for this and other diseases [28–30]. Although various plants have been traditionally used throughout history to reduce blood glucose and improve diabetes complications, there is not enough scientific information about some of them. Herbal medicines are commonly prescribed throughout the world because of low side effects, availability, roughly low cost, and also its effectiveness [31, 32].

In Saudi Arabia, the number of people who suffer from DM increased from 890,000 in 2000 to a staggering projection of 2,523,000 in 2030. In 2011, Saudi Arabia reported a prevalence of DM at 30% of the total population, with a rate of 27.6% in women and 34.1% in men [33]. According to 2010 data from several sources (WHO, World Bank, UNESCO, CIA, and individual country databases), DM is the number three disease-related cause of death in Saudi Arabia [34].

In the present situation, herbal medicines’ usage has significantly increased and published studies from developed countries emphasize that a paramount proportion of medicines supplied by them have herbal origins, so growing and producing the herbal medicines could be helpful to both economic development and community’s health [35]. Keeping in mind the importance of medicinal plants, in the current review various medicinal plants used for antidiabetic treatment around the world, native to or cultivated in Saudi Arabia, are documented for the purpose to provide up-to-date insight on medicinal plant used for DM, so that researcher easily selects plant for bioscreening and active constituents’ identification purposes. Therefore, we invite researchers’ attention to carry out detailed ethnopharmacological and toxicological studies on unexplored antidiabetic plants in order to provide reliable knowledge to the patients and develop novel antidiabetic drugs.

2. Methods

Required data about medicinal plants having antidiabetic activities and growing in the Kingdom of Saudi Arabia were searched/collected from the online databases including Wiley, Google, PubMed, Google Scholar, ScienceDirect, and Scopus. Keywords used in search are in vivo antidiabetic activities, flora of Saudi Arabia, active ingredients, toxicological evaluations, and medicinal plants. Latest published data approximately in the last ten years with the key outcome of change in blood glucose level in animal model were included. One or two articles are selected as references for each plant’s species on priority basis from the journals found in web of science and latest years.

3. Results

The names, families, used parts, location, and antidiabetic properties in animal model of the native/cultivated Saudi medicinal plants are summarized in Table 1. The active ingredients and toxicological effect of these plants in animal model are given in Table 2. A total of 50 plant species belong from 27 families were found in the flora of Saudi Arabia. Dominant family was found Lamiaceae with 5 species (highest) followed by Moraceae with 4 species.

4. Discussion

The majority of the experiments confirmed the benefits of medicinal plants with hypoglycemic effects in the management of diabetes mellitus. From Table 1, it can be concluded that among the plants used for the treatment of diabetes, H. salicornicum, T. oliverianum, A. cepa, A. herba-alba, Teucrium polium, Sesamum indicum, Z. spina-christi, and U. dioica seem to be most common plants used to treat diabetes and are available everywhere in the world. The leaves were most commonly used plant part, and other parts (root, stem, bark, flower, seed, and whole plant) were also useful for curing. The most common diabetic model that was used was the streptozotocin and alloxan-induced diabetic mouse or rat as diabetic models. The most commonly involved active constituents are flavonoid, alkaloid, saponin, carbohydrate, vitamins, amino acid and its derivatives, phenol and its derivatives, and benzoic acid derivatives. The very common phytoconstituents, targeted metabolic pathways, and its structure are given in Table 3 [194, 195]. The native to or cultivated plant species of the kingdom given in Table 1 are selected from the published literature about ethnobotanical value and antidiabetic potential of medicinal
| S. no. | Names of plants | Family | Part used | location | Antidiabetes Activities |
|-------|-----------------|--------|-----------|----------|------------------------|
| 1.    | *Allium cepa*   | Liliaceae | Bulb      | Central Saudi Arabia [36] | Ethanol extract of *A. cepa* in STZ-induced diabetic rats causes 66% decreased at 200 mg/kg after 24 h in blood glucose level [37]. 0.4 g/100 g bw of *A. cepa* reduced 50% the fasting glucose levels of diabetic rats [38]. Similar results reported by other researchers [39]. |
| 2.    | *Anthemis herba-alba* | Compositae/ Asteraceae | Aerial parts | Farasan Island of Red Sea [40] | 72% plasma glucose levels decreased in albino mice by ethyl alcohol extract of *Artemisia herba-alba* [41]. *C. intybus* leaf powder, ethanol, aqueous seed extracts, and hexane extracts led to a decrease in blood glucose levels to near normal value. |
| 3.    | *Cichorium intybus* | Asteraceae | Seeds | Qassim region [42] | Hypoglycemic effects of *C. intybus* were observed in diabetic rats, and a dose of 125 mg of plant extract/kg body weight exhibited the most potent hypoglycemic effect [43–45]. |
| 4.    | *Clitoria ternatea* | Fabaceae | Aerial parts | Cultivated throughout Saudi Arabia [46] | The aqueous extract of *Clitoria ternatea* leaves and flower administered for 84 days to diabetic rats significantly decreased blood glucose [46–48]. Different extracts and fractions of *F. carica* showed a clear hypoglycemic effect in diabetic rats. |
| 5.    | *Ficus carica* | Moraceae | Leaves | Southwest of Saudi Arabia [49] | *F. carica* leaves exerted significant effect on carbohydrate metabolism enzymes with promising hypoglycemic and hypolipidemic activities in type 2 diabetic rats [50, 51]. In streptozotocin-induced diabetic rats, bark aqueous extract, and an isolated compound, α-amyrin acetate exhibited antidiabetic activity by decreasing the blood glucose level and increasing the HDL level [53]. |
| 6.    | *Ficus benghalensis* | Moraceae | Bark | Riyadh [52] | The aqueous extract of bark and ethanol extract of leaves and fruits had a promising antidiabetic effect in streptozotocin-induced diabetic rats by decreasing the blood glucose, serum triglyceride, and total cholesterol levels and increasing serum insulin, body weight, and glycogen content in the liver and skeletal muscle [53]. |
| 7.    | *Ficus religiosa* | Moraceae | Root bark, stem bark, aerial roots | Riyadh [52] | *F. microcarpa* leaves showed protective effect against alloxan-induced diabetic rats by reducing blood glucose, cholesterol and triglyceride levels, and increased insulin level [53]. |
| 8.    | *Ficus microcarpa* | Moraceae | Leaves | Riyadh [52] | H. perforatum ethyl acetate extract possesses potent antihyperglycemic activity in STZ-induced diabetic rats [55]. |
| 9.    | *Hypericum perforatum* | Hypericaceae | Leaves | Western Saudi Arabia [54] | Different extracts and tablets of *Anethum graveolens* possess potent antihyperglycemic activity in alloxan-induced diabetic mice [57]. Oral administration of cumin seeds crude ethanol extract and glibenclamide to diabetic rats significantly and progressively restored toward normal. Cumin seeds crude ethanol extract and glibenclamide reduced plasma glucose levels by 38.34 and 37.73%, respectively, compared with diabetic control [58]. Other studies also reported similar results [59]. |
| 10.   | *Anethum graveolens* | Apiaceae | Seeds | Makka [56] | M. vulgar extracts lower blood glucose level 30 to 60% in dose-dependent manner in streptozotocin-induced diabetic rats [60]. |
| 11.   | *Cuminum cyminum L.* | Apiaceae or Umbelliferae | Seeds | Makka [56] | |
| S. no. | Names of plants      | Family       | Part used | location         | Antidiabetes Activities                                                                 |
|-------|----------------------|--------------|-----------|------------------|-----------------------------------------------------------------------------------------|
| 13    | Mentha longifolia    | Lamiaceae    | Whole plant | Madinah [61]     | Remarkable antidiabetic, anticholinesterase, and antityrosinase effects were recorded for the mint oil [61, 62]. Still need to investigate in vivo antidiabetic potential. |
| 14    | Origanum syriacum    | Lamiaceae    | Leaves    | Saudi Desert [63] | The whole plant extract of O. syriacum at 100 and 400 mg/kg significantly lowers glucose level in diabetic induced rats [64]. Aqueous and ethanol extract of Teucrium oliverianum were tested for antidiabetic activity in alloxan-induced diabetic mice. Both extracts significantly reduced blood sugar levels [65] Infusion orally (64% decrease glucose level) and intraperitoneal of different extracts of T. polium caused significant reductions in blood glucose concentration in STZ hyperglycemic rats [68] |
| 15    | Teucrium oliverianum | Lamiaceae    | Aerial parts | Throughout Saudi Arabia [65,66] |                                                                                           |
| 16    | Teucrium polium      | Lamiaceae    | Leaves    | Madinah [67]     |                                                                                           |
| 17    | Achyranthes aspera   | Amaranthaceae | Whole plant | Al Hada Road Taif [69] |                                                                                           |
| 18    | Aerva lanata         | Amaranthaceae | Leaves    | Southwest region of Saudi Arabia [71, 72] |                                                                                           |
| 19    | Alternanthera sessilis | Amaranthaceae | Whole plant | Hail region, Saudi Arabia [74] |                                                                                           |
| 20    | Carissa edulis       | Apocynaceae  | Leaves    | Southern region of Saudi Arabia [76] | Oral administration of C. edulis extracts of the leaves significantly reduced the blood glucose level in STZ diabetic rats [77]. C. roseus (100 mg/kg BW) lowered the glucose level more than metformin-treated group (100 mg/kg BW) in STZ-induced hyperglycemia rats. C. roseus 200 mg/kg dose was found to be more effective in reducing fasting blood glucose levels [79] |
| 21    | Catharanthus roseus  | Apocynaceae  | Flower, leaves, stem, and root | Western Saudi Arabia [78] | Extracts Rhazya stricta lowered 37.9% blood glucose level in the streptozotocin-induced diabetic rats. Serum cholesterol and triglyceride levels were significantly (P < 0.05) reduced in the treated diabetic group compared to the untreated diabetic group [81] Different extracts of C. procera at dose of 250 mg/kg were orally administered as single dose per day to diabetes-induced rats for the period of 15 days significantly decreases blood glucose level to the level of standard drug glibenclamide [83] Researcher observed the significant hypoglycemic activity of Opuntia dillenii extract in streptozotocin-induced diabetic mice and rabbits [85] |
| 22    | Rhaiza stricta       | Apocynaceae  | Leaves, seeds | Middle and western region of Saudi Arabia [80] |                                                                                           |
| 23    | Calotropis procera   | Asclepiadaceae | Latex    | Al-Kharj [82]     |                                                                                           |
| 24    | Opuntia dillenii     | Cactaceae    | Fruit     | Jazan Region [84] |                                                                                           |
| S. no. | Names of plants         | Family           | Part used   | location                      | Antidiabetes Activities                                                                                                                                                                                                 |
|-------|------------------------|------------------|-------------|-------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 25    | *Opuntia ficus-indica* | Cactaceae        | Stem        | Jazan Region [84]             | Powder and water extract of *O. ficus-indica* significantly (in comparison with control group) returned blood glucose level to the initial level, 180 min after administration in STZ-induced diabetic rats [86]. Many studies confirmed the hypoglycemic activities of *O. ficus-indica* [87]. C. *decidua* extracts at dose level of 200 and 800 mg/kg significantly reduce sugar level (in a dose-dependent manner) compared to standard drug in STZ-induced diabetic and normal rats [88]. Extract of *B. vulgaris* at does level 50, 100, and 200 mg/kg of significantly reduced sugar level and increased in insulin level (in a dose-dependent manner) in streptozotocin or alloxan-induced diabetic mice [90]. Other researchers also concluded similar finding in STZ-induced diabetic rats [91]. |
| 26    | *Capparis decidua*     | Capparaceae      | Fruits, seeds | Jazan Region [84]             | C. *decidua* extract at dose level 200 and 800 mg/kg significantly reduce sugar level (in a dose-dependent manner) compared to standard drug in STZ-induced diabetic and normal rats [88]. Extract of *B. vulgaris* at does level 50, 100, and 200 mg/kg of significantly reduced sugar level and increased in insulin level (in a dose-dependent manner) in streptozotocin or alloxan-induced diabetic mice [90]. Other researchers also concluded similar finding in STZ-induced diabetic rats [91]. |
| 27    | *Beta vulgaris*        | Chenopodiaceae   | Root bark   | North Hejaz and Eastern Najd region of Saudi Arabia [89] | Ethanol extract (100 and 200 mg/kg of bw) of *H. salicornium* (oral administration) exhibited persistent hypoglycemic effects in STZ-induced diabetic rats [93]. *E. alsinoides* ethanol extract at dose level (150 mg/kg bw) in normal and streptozotocin-induced diabetic rats leads to hyperglycemia in experimental diabetic rats that decreased utilization of glucose by insulin-dependent pathways [94, 95]. |
| 28    | *Haloxylon salicornicum* Bunge | Chenopodiaceae | Whole plant | Wadi-Hafir-Al-Batin, Saudi Arabia [92] | *I. aquatica* ethanol extract at dose level (10, 100, and 1000 µg/ml in streptozotocin-induced diabetic rats significantly (P < .05) exhibited the ability to enhance insulin-mediated glucose uptake into 3T3-F442A adipocytes cells compared to insulin alone [97]. Another study confirmed that doses (200 mg/kg and 400 mg/kg) reduced blood glucose level, and it was statistically highly significant (P < 0.001) in comparison with control group [98]. 1 ml/kg and 2 ml/kg of *C. colocynthis* extract (orally administered) stabilized animal body weight and ameliorated hyperglycemia in a dose- and time-dependent manner in alloxan-induced diabetic rats [99]. |
| 29    | *Evolvulus alsinoides*  | Convolvulaceae   | Whole plant | Jazan Region [84]             | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| 30    | *Ipomea aquatica*      | Convolvulaceae   | Whole plant | Jazan Region [96]             | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| 31    | *Citrullus colocynthis* | Cucurbitaceae   | Fruits      | Jazan Region [84]             | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| 32    | *Citrullus lanatus*     | Cucurbitaceae   | Seed        | Wadi Lajab, Saudi Arabia [100] | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| 33    | *Coccinia grandis*     | Cucurbitaceae   | Whole plant | Jazan Region [84]             | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| 34    | *Jatropha curcas*       | Euphorbiaceae   | Leaves      | Jazan Region [84]             | *C. colocynthis* seed extract (2, 4 g/kg) treatment significantly lowers glucose level which suggested that *C. lanatus* had antidiabetic property in STZ-induced diabetes mice [101]. Other researcher also concluded similar finding in STZ-induced diabetic rats [102]. The C. *grandis* extract (0.75 mg/kg, orally) showed remarkable glycemic effect which confirmed antidiabetic potential in streptozotocin-induced diabetic rats [103]. Ethanol extract of *J. curcas* leaves at doses of (250 and 500 mg ml⁻¹ bw by administered orally) reduced glucose level from 219.5 to 116.5 and 237 to 98.8, respectively, in alloxan-induced diabetic rats. The results were comparable to reduction in rats treated with the standard glibenclamide 232–94.5 at 600 µg kg⁻¹ [104]. |
| S. no. | Names of plants     | Family          | Part used | location           | Antidiabetes Activities                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
|-------|--------------------|-----------------|-----------|--------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 35    | *Ricinus communis* | Euphorbiaceae   | Leaves    | Jazan Region [84]  | *R. communis* extracts at doses of 300 and 600 mg/kg/BW administered orally caused hyperglycemia in a dose-dependent manner in streptozotocin-induced diabetic rats [105]. A review article focusing on antidiabetic potential of *F. carica* confirmed that different extracts and fractions of *F. carica* and different doses significantly reducing hyperglycemia in streptozotocin-induced diabetic rats compared to standard drug [106]. Alloxan-induced type 2 diabetic albino Wistar rats treated with 250, 500, and 1000 mg/kg (body weight) of the extract of *F. sycomorus* intraperitoneally reduced glucose level in diabetic rats almost to the normal as compared to diabetic control [107]. Alloxan-induced diabetic rats treated with 5% and 10% of *Sesamum indicum* seed powder significantly decreased blood glucose and increased insulin levels as compared with the positive (diabetic) control group [108]. In intravenous administration of alloxan-induced diabetic rabbits glucose level lowering effect observed (time dependent manner) with *P. ovata* husk extract of dose level (300 mg/kg, orally administered) [109]. 0.7 g/kg of *P. erioptera* extract showed significant antidiabetic effect compared to standard drug metformin and glibenclamide in normal and alloxan-induced diabetic rats [110]. Many ethnopharmacological investigations reported its antidiabetic potential but still need to study its in vivo and in vitro antidiabetic potential [113, 114]. The strongest (*P* < 0.001) antidiabetic activity (25.59 and 39.48% after 7 and 15 days, respectively) was found following treatment with dose level of 500 mg/kg of *Z. spina-christi* extract in streptozotocin-induced diabetes mice [115]. *B. monnieri* extract at dose level of 50, 100, 200, and 400 mg/kg significantly inhibited (33.3, 34.2, 42.1, and 44.2%, respectively) the increase in serum glucose concentration in a dose-dependent manner compared to standard drug [116]. The strongest (*P* < 0.001) antidiabetic activity of *L. shawii* extract of 250 and 500 mg/kg bw was found in a dose-dependent manner in streptozotocin-induced diabetes rats [117]. *S. nigrum* extract was given orally in the dose level of 200 and 400 mg/kg/day (7 days) significantly lowering the blood glucose level in fasting compared to standard drug in alloxan-induced diabetic albino Wistar rats [118]. |
| 36    | *Ficus carica*      | Moraceae        | Leaves    | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 37    | *Ficus sycomorus*   | Moraceae        | Leaves    | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 38    | *Sesamum indicum*   | Pedaliaceae     | Seeds     | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 39    | *Plantago ovata*    | Plantaginaceae  | Husk      | Northern border region of Saudi Arabia [18] |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 40    | *Polygala erioptera*| Polygalaceae    | Aerial part | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 41    | *Polygonum aviculare* L | Polygonaceae | Aerial parts | Taif Region [111, 112] |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 42    | *Ziziphus spina-christi* | Rhamnaceae | Leaves    | Eastern region of Saudi Arabia [84, 112] |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 43    | *Bacopa monnieri*   | Scrophulariaceae| Aerial parts | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 44    | *Lycium shawii*     | Solanaceae      | Aerial parts | Taif Region [112]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 45    | *Solanum nigrum*    | Solanaceae      | Whole plant | Jazan Region [84]  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
plants around the world. The ethnobotanical information reports about 800 plants that may possess antidiabetic potential [196, 197]. Jeeva and Anlin also reported 177 plants belonging to 156 genera and 76 families used traditionally for antidiabetic treatment [198]. In the Middle East countries, there are 129 plant species still in use in traditional Arabic medicine. This indicates that the medicinal plant species require preservation as well as the ethnobotanical and ethnopharmacological knowledge. The preservation of the herbs is an essential requirement for maintaining traditional Arabic medicine as a medicinal and cultural resource [199]. The selected plant species *H. salicornicum*, *T. oliverianum*, *A. cepa*, *A. herba-alba*, *Teucrium polium*, *Sesamum indicum*, *Z. spina-christi*, and *F. religiosa* are the native Saudi medicinal plants traditionally used for the treatment of DM [200]. Similarly published data showed that 20 medicinal plants are traditionally used in Tabuk region of Saudi Arabia [201]. *Anisotes trilicus*, *Artemisia judaica*, and *Moringa peregrine* are used in Al Khobah village, Saudi Arabia, for DM treatment [202]. *O. europaea* is used in Al Bahah region of KSA for DM treatment [203]. *C. roseus*, *A. cepa*, *U. dioica*, *A. aspera*, *C. intybus*, *C. cyminum*, *F. bengalensis*, *C. colocynthis*, and *T. polium* are the highly investigated medicinal plants for antidiabetic potential [204–206].

Desiring to contribute to the conservation priorities of traditional medicine knowledge of various medicinal plants native to or cultivated in Saudi Arabia and to make it easy and familiarized with disease treatment, the present compilation was conducted. According to the International Union for Conservation of Nature and the World Wildlife Fund, there about 15,000 medicinal plant species are threatened with extinction from overharvesting and habitat destruction and 20% of their wild resources have already been nearly exhausted with the increasing human population and plant consumption [207]. Each plant species lost due to extinction phenomena could represent not only the loss of healthcare saving cures for special diseases but also the loss of probable primary metabolite like protein- or vitamin-rich foods [208]. Medicinal plants have been cited as a potential source of heavy metal toxicity to both man and animals. The most common heavy metals implicated in human toxicity include lead, mercury, arsenic, and cadmium, although aluminum and cobalt may also cause

| S. no. | Names of plants | Family | Part used | location | Antidiabetes Activities |
|-------|----------------|--------|-----------|----------|-------------------------|
| 46    | *Withania somnifera* | Solanaceae | Leaves | Jazan Region [84] | *W. somnifera* extract oral administration at two doses (200 and 400 mg/kg) reduced the blood glucose level significantly (*P* < 0.001) in a dose-dependent manner in streptozotocin-induced diabetes rats. Only WS treatment did not register any significant change in the blood glucose level when compared to citrate control rats [119]. Another study also confirmed similar results in alloxan-induced diabetic rats [120]. Literature survey showed that *L. camara* leaf extract oral administration (200, 250, and 500 mg/kg of bw) showed antidiabetic potential in alloxan-induced diabetic rats [121]. *P. harmala* seed extract at dose level of (30, 60, and 120 mg/kg, orally administered for four weeks) significantly decreases in blood glucose (in all doses, *P* < 0.001), in comparison with diabetic group [122]. *T. terrestris* extract at (2 g/kg body weight) produced protective effect in streptozotocin-induced diabetic rats by inhibiting oxidative stress [123]. *T. terrestris* L. extract (250 mg/kg of bw orally administered) significantly lowers glucose level to normal compared to standard drug in glucose-loaded normal rabbits [124]. *Urtica dioica* extract at 100 mg/kg (*P* < 0.01) and 200 mg/kg (*P* < 0.001) significantly decreased serum glucose fructose-induced insulin resistance rats [126]. The aqueous extract of *U. dioica* significantly (*P* < 0.001; 67.92%) reduced the blood glucose level at dose of 300 mg/kg, IP in streptozotocin-induced diabetes rats [127]. |
| 47    | *Lantana camara* | Verbenaceae | Leaves | Jazan Region [84] |
| 48    | *Peganum harmala* | Zygophyllaceae | Seeds | Taif Region [112] |
| 49    | *Tribulus terrestris* | Zygophyllaceae | Stem, leaves | Jazan Region [84] | Taif Region [112] |
| 50    | *Urtica dioica* | Urticaceae | Leaves | Wild plant, Tanhat, Saudi Arabia [125] |
Table 2: Active ingredients and toxicological evaluation of the medicinal plants given in Table 1.

| S. No | Names                  | Active ingredients                                                                 | Toxicological evaluation                                                                 |
|-------|------------------------|-------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| 1     | Allium cepa            | Quercetin, N-acetylcysteine, alliucioide, cycloalliin, S-methyl-L-cysteine, S-propyl-L-cysteine, sulfoxide, dimethyltrisulfide, S-methyl-L-cysteine sulfoxide [128] | The animals tested were found healthy with no sign of toxicity up to the dose of 2,500 mg/kg. However, at 5,000 mg/kg, animals were weak and had intense extreme tachycardia and disorientation but no death was recorded. Thus, LD<sub>50</sub> was more than 5,000 mg/kg [129]. |
| 2     | Anthemis herba-alba     | Guainalides, eudesmanolide, pseudouga inolides, xanthonolides, flavone, flavonol glycosides, hispidulin, cirsilinole, vicenin-2, safrasoids, isoschaftsoides, 5',4'-dihydroxy-6,7,3-trimethoxyflavone, quercetin-3-rutinoside, patuletin 3-rutinoside, patuletin 3-glucoside [130] | The available toxicological investigations have shown generally that Anthemis herba-alba is free from toxic effects at the different doses used in the studies [130] |
| 3     | Cichorium intybus       | Chicoric acid, inulin, cichoralexin, cichorin, esculetin, isochlorogenic acid, chlorogenic acid, caffeic acid, dicafeoylquinic acid, aesculin, arginine, histidine, isoleucine, leucine, lysine, methionine, cysteine, phyllanalnine, tyrosine, threonine, valine, serine, glutamic acid, glycine, alanine, asparatic acid, and proline [44, 45] | Ethanolic extract of aerial parts and root of CT led to lethargy in mice at the doses of 1500 mg/kg and above, orally [31]. Proxit was seen above 2000 mg/kg dose in mice. Through intraperitoneal route, 2900 mg/kg dose was lethal within 6 hr due to severe CNS depression [47]. The rats tested were found healthy with no sign of toxicity up to the dose of 5000, 5500, and 6000 mg/kg. However, at 5000 mg/kg, animals were weak and had intense extreme tachycardia and disorientation but no death was recorded. Thus, LD<sub>50</sub> was more than 6000 mg/kg [132]. |
| 4     | Clitoria ternatea       | Kaempferol, quercetin, myricetin, taxaxerol, tannic acid, 3-monoglucoside, β-sitosterol, delphinidin-3,5-diglucoside, anthoxanthin glucoside, p-hydroxycinnamic acid, kaempferol 3-neohesperidoside, myricetin 3-rutinoside, hexacosanol [48] | There were no treatment-related toxic effects from chicory extract administered orally at 70, 350, or 1000 mg/kg/day. There were no observed adverse effects of chicory extract in these studies [45] |
| 5     | Ficus carica            | Over 100 bioactive compounds have been identified in fig such as rutin, arabinose, chlorogenic acid, β-amyrins, syringic acid, β-carotenones, glycosides, β-sitosterols, and xanthotinol [131] | In acute toxicity studies, no mortality and signs of toxicity were observed at the dose of 2000 and 5000 mg/kg body weight for aqueous and ethanol extracts, respectively [53] |
| 6     | Ficus benghalensis      | Leucopelargonidin-3-0-α-L. rhamnoside, eucodelphinidin, leucoanthocyanidin, leucoanthocyanin, α-amyrin acetate [53] | Acute toxicity reported up to dose level 2000 mg/kg showed no mortality [53] |
| 7     | Ficus religiosa         | Lupeol, β-sitosterol, β-sitosterol-d-glucoside, stigmasterol, lanosterol, campesterol, octacosanol, methyl oleonate, lupen-3-one, bergapten, and bergaptole [53] | The oral administration of a single dose of 2000 mg/kg ethanol or methanol extract of leaves showed no mortality or behavioral alterations in the tested animals [53] |
| 8     | Ficus microcarpa        | Polyphenols, organic acids, alkaloids, polysaccharides, megastigmanes, pheophytins, catechin, epicatechin, isovitexin, phenolic acids [53] | Acute toxicity studies revealed the nontoxic nature of the H. perforatum [55]. The mice treated with AG of different doses of 1000, 2000, 3000, 4000, and 5000 mg/kg of body showed no toxicity [135] |
| 9     | Hypericum perforatum    | Quercitrin, rutin, hypericin, kaempferol, biapigenin, hyperforin [133] | The acute lethal toxicity test revealed that cumin crude extract was very safe [58]. An acute toxicity study of M. vulgare (1 g/kg) extract orally administered at a dose of 1 g/kg body weight to the mice and treated mice showed tachycardia 1 h after intake of the infusion and loss of appetite 3 h after intake of the infusion. In another experiment, a single dose of 2000 mg/kg extract of M. vulgare for an acute toxicity study showed no toxicity [60]. |
| 10    | Anethum graveolens      | Carvone, α-phellandrene, limonene, dill ether, myristicin coumarins, flavonoids, phenolic acids, steroids [134] | |
| 11    | Cuminum cyminum         | Cuminaldehyde, limonene, α- and β-pinene, 1, 8-cineole, α- and p-cymene, α- and γ-terpinene, safranal, and linalool [58, 59] | |
| 12    | Marrubium vulgare       | Furanic labdane diterpenes, marrubiol, marrubiin, ladanein [60] | |
| S. No | Names                   | Active ingredients                                                                                                                                                                                                 | Toxicological evaluation                                                                                                                                                                                                 |
|-------|-------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 13    | *Mentha longifolia*     | Mentha-1, mentha-2, camphelione, camphene, carveol, carvone, carvone oxide, lirionene, linalool, menthatriene, menthofuran, menthol, menthone, myrcene, p-cymene, piperitene, pipertone, piperitone, sabinene, α-pinene, α-terpinene, α-terpineol, longifone, pulegone, longifolione-A, longifolione-B, longiside-B, eugenol, salvianolic acid, eriodictyol-7-rutinoside, apigenin-7-O-glucoside, hypolaetin, longitin, luteolin, etc. [136] | *M. longifolia* extract was safe, and no toxicity or mortality was observed in both the oral (3200 mg/kg) and intraperitoneal (1730 mg/kg) administration in rats. Fourteen days of oral administration of the essential oil (125, 250, 375, and 500 µL/kg) resulted in the reduction of red blood cells and lymphocytes and elevation of neutrophils and monocytes compared with normal animals [136]. |
| 14    | *Origanum syriacum*     | Carvacrol, thymol, thymoquinone [137]                                                                                                                                                                                  | Not available                                                                                                                                                                                                          |
| 15    | *Teucrium oliverianum*  | 8-O-acetylharpagide, 12-O-methylteucrolin A, teucrolivin A, eupatorin, teucrolivin B, μ24(S)-stigmasta-5,22,25-trin-3β-ol [66]                                                                                       | Not available                                                                                                                                                                                                          |
| 16    | *Teucrium polium*       | Apigenin, luteolin, rutin, cirsimaritin, salvigenin, and eupatorin in the roots, aerial parts, and inflorescences, teucardioside, b-sitosterol, stigmasterol, campesterol, brassicasterol, and clerosterol [68] | All rats treated with different concentrations of the total extract of TP were alive during the 14 days of observation. The animals did not show visible signs of acute toxicity. It suggested that the LD50 of the total extract was higher than 8 g/kg [138]. |
| 17    | *Achyranthes aspera*    | Aliphatic acid, betaene, achyranthine, β-ecdysterone, achyranthes saponins A, B, C, D, oleononic acid, glycosides, triacanotal, E-sitosterol and spinasterol, triacanotal, hydroquinone, eugenol [70] | In acute oral toxicity studies, there was no increase or decrease in any of the parameters studied, in comparison with control animals [139]. The LD50 of the extract of AL for oral and IP acute toxicity tests were 22.62 g/kg and 0.432 g/kg, respectively. The extract produced apparent changes in body weights of both male and female rats and increased the weights of lung, brain, and pancreas of female rats while reducing the weight of testes in male rats. Hematological parameters were also altered [72]. |
| 18    | *Aerva lanata*          | Quercetin, betulin, aevrione, eroside, methylervbine, aevrione, lupeol, kaempferol, aevrolamine, aervoline, eroside, METHYLAERVINE, PERSINOSIDES A AND B, TANNIC ACID, LUPEOL ACETATE, BENZOIC ACID, METHYL GREVILLATE [140] | TheLD50 of the crude extract did not show any toxicity in mice even at the highest dose tested [75]. Lethal effects were not observed after the oral administration of the standardized ethanol extract at doses of 1600, 2900, and 5000 mg/kg. No behavioral changes were observed during the observation period. The oral LD50 of the extract was estimated to be greater than 5000 mg/kg. [142]. No mortality, but dose level higher than 300 mg of *C. roseus* extract can produce signs of biochemical and histopathological toxicity in liver, kidney, and heart. It is recommended that lower doses than the studied ones should be used for treatment [144]. |
| 19    | *Alternanthera sessilis* | Stigmasterol, β-sitosterol, β-carotene, ricinoleic acid, myristic, palmitic, stearic, oleic, and linoleic acids, α-spiraterol, uronic acid, cyclo-eucalenol, choline, oleanolic acid, lupeol [141] | The crude extract did not show any toxicity in mice even at the highest dose tested [75]. Lethal effects were not observed after the oral administration of the standardized ethanol extract at doses of 1600, 2900, and 5000 mg/kg. No behavioral changes were observed during the observation period. The oral LD50 of the extract was estimated to be greater than 5000 mg/kg. [142]. No mortality, but dose level higher than 300 mg of *C. roseus* extract can produce signs of biochemical and histopathological toxicity in liver, kidney, and heart. It is recommended that lower doses than the studied ones should be used for treatment [144]. |
| 20    | *Carissa edulis*        | B-Amyrin, (+)-carissone, 2α-carissanol, 6α-carissanol, dehydrocarissone, pinene, myrcene, limonene, subanene, rutin, epicatechin gallate, carinol, lariresinol, β-sitosterol, sitosterol glucoside, stigmasterol glucoside, scopoletin, isofoxaxadin, pinitol [77] | The crude extract did not show any toxicity in mice even at the highest dose tested [75]. Lethal effects were not observed after the oral administration of the standardized ethanol extract at doses of 1600, 2900, and 5000 mg/kg. No behavioral changes were observed during the observation period. The oral LD50 of the extract was estimated to be greater than 5000 mg/kg. [142]. No mortality, but dose level higher than 300 mg of *C. roseus* extract can produce signs of biochemical and histopathological toxicity in liver, kidney, and heart. It is recommended that lower doses than the studied ones should be used for treatment [144]. |
| 21    | *Catharanthus roseus*   | Vinblastine, vincristine, vindesine, vindoline tabersonine, ajmalicine, vinceine, vineamine, raubasin, reserpine, catharanthine, rosinind [143]                                                                          | Daily oral dosing of *R. stricta* extract (0.25 g/kg) for 42 days was not fatal to sheep [145]. |
| 22    | *Rhazya stricta*        | Polyneuridine, stemmadenine, strictanol, rhazimine, rhazinilam, rhazimamine, sewarine, vallesiacotamine, tetrahydrosecomeine [145]                                                                               | Daily oral dosing of *R. stricta* extract (0.25 g/kg) for 42 days was not fatal to sheep [145]. |
| S. No | Names                         | Active ingredients                                                                 | Toxicological evaluation                                                                                                                                 |
|-------|-------------------------------|-------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------|
| 23    | Calotropis procera            | Betalin, betanidin, kaempferol, kaempferide, quercetin, isorhamnetin, β-sitosterol, | 2000 mg/kg body weight in single oral administration of aqueous and hydroalcoholic extract did not cause any death after 72 h post-treatment in male and female mice. Daily administration of aqueous extract to male and female Wistar rats during 3 and 6 weeks at the dose of 20 mg/kg/day induced no mortality in either sex [147]. Whoever C. procera is a toxic plant that is avoided by grazing animals. Its latex is used by tribes to poison arrows used for hunting. If in contact with human eye, it could cause ocular toxicity, causing loss of vision and photophobia [146] |
|       |                               | C29-5β-sterols, taraxerol, methyl linoleate, 7-oxosterol, 6β-hydroxystigmast-4-ene-3-one, daucosterol, methyl eucomate, eucomic acid [85] |                                                                                                                                                          |
| 24    | Opuntia dillenii              | Quercetin, isorhamnetin, kaempferol, luteolin, n-triacontane, n-pentacosane, β-carotene, n-triacontanol, kaempferol, quercetin, isodulcite, nanocosane, capric acid, glucocapparin, capparine, capparinine, cappariline, codonocarpin, β-sitosterol [88] | During the oral toxicity study of the crude drug in rats, given doses up to 50 ml/kg exhibited no symptoms of toxicity [85] |
| 25    | Opuntia ficus indica          | Isorhamnetin glycosides, gallic acid, coumaric, nitrin, totarol, isoquercetin, ferulic acid [87], C. colocynthis extract did not cause any mortality or adverse effect [92]. | In vivo toxicity study suggests that the oral administration of Opuntia ficus indica extract at levels up to 2000 mg/kg/day does not cause adverse effects in male and female rats [148]. The oral administration of C. deciduala extract (500, 1000, 2000, and 4000 mg/kg) did not provoke any gross behavioral changes or manifestations of toxic symptoms in male rats [149]. In acute oral toxicity studies, the BVBF did not show any sign and symptoms of toxicity and mortality up to 2000 mg/kg dose, considered relatively safe [150] |
| 26    | Capparis decidua              | Kaempferol, quercetin, betaine chloride, piperidine, anabasine, aldotripterideine, haloxine, halosaline, oxeidine, tyramine, N-methyltyramine, scopoletin, scopolin, umbelliferone, xanthotoxol, isoxyperatorin, esculin, β-sitosterol, ursolic acid, β-amyrin [93], β-Sitosterol, betaine, shankpushpin, evolvine, caffeic acid, 6-methoxy-7-O-β-glucopyranoside coumarin, 2-C-methyl erythritol, kaempferol-7-O-β-glucopyranoside, kaempferol-3-O-β-glucopyranoside, quercetin-3-O-β-glucopyranoside, scopoletin, scopolin [151] Caffeic acid, chlorogenic acid, quercetin glucoside, quercetin malonyl glucoside, quercetin diglucoside, catechin, isochlorogenic acid A, C, aspartic acid, glycine, alanine and leucine, 7-O-β-D-glucopyranosyldihydroquercetin-3-O-α-D-glucopyranoside [97, 98] | The Evolvlus alsinoides extract did not cause any mortality up to a dose of 1500 mg/kg body weight and no behavioral, neurological, and autonomic profiles and was found to be safe [151]. |
| 27    | Beta vulgaris                 | Betaine, betacyanins, betaxanthins, oxalic acid, and ascorbic acid [89]            | H. salicorlicum extract at doses 0.1, 0.2, 0.3, 0.4, and 0.5 mL/kg orally administered in rats was safe and showed no mortality or adverse effect [92]. |
| 28    | Haloxylon salicorlicum        | Kaempferol, quercetin, betaine chloride, piperidine, anabasine, aldotripterideine, haloxine, halosaline, oxeidine, tyramine, N-methyltyramine, scopoletin, scopolin, umbelliferone, xanthotoxol, isoxyperatorin, esculentin, β-sitosterol, ursolic acid, β-amyrin [93]. | The Evolvlus alsinoides extract did not cause any mortality up to a dose of 1500 mg/kg body weight and no behavioral, neurological, and autonomic profiles and was found to be safe [151]. |
|       | Bunge                        | β-Sitosterol, betaine, shankpushpin, evolvine, caffeic acid, 6-methoxy-7-O-β-glucopyranoside coumarin, 2-C-methyl erythritol, kaempferol-7-O-β-glucopyranoside, kaempferol-3-O-β-glucopyranoside, quercetin-3-O-β-glucopyranoside, scopoletin, scopolin [151] Caffeic acid, chlorogenic acid, quercetin glucoside, quercetin malonyl glucoside, quercetin diglucoside, catechin, isochlorogenic acid A, C, aspartic acid, glycine, alanine and leucine, 7-O-β-D-glucopyranosyldihydroquercetin-3-O-α-D-glucopyranoside [97, 98] | The Evolvlus alsinoides extract did not cause any mortality up to a dose of 1500 mg/kg body weight and no behavioral, neurological, and autonomic profiles and was found to be safe [151]. |
| 29    | Evolvulus alsinoides          | Caffeic acid, chlorogenic acid, quercetin glucoside, quercetin malonyl glucoside, quercetin diglucoside, catechin, isochlorogenic acid A, C, aspartic acid, glycine, alanine and leucine, 7-O-β-D-glucopyranosyldihydroquercetin-3-O-α-D-glucopyranoside [97, 98] | The Evolvlus alsinoides extract did not cause any mortality up to a dose of 1500 mg/kg body weight and no behavioral, neurological, and autonomic profiles and was found to be safe [151]. |
| 30    | Ipomea aquatica              | Cucurbitane, gallic acid, kaempferol, cucurbitacin A-E, I-L, chlorogenic acid, caffeic acid, colochinosides A, B, C; choline, almitic acid, stearic acid, linoleic acid, oleic acids, catechin, myricetin, α-tocopherol, y-tocopherol, β-carotene [153] | In acute toxicity studies, I. aquatica extract was found to be safe up to 2g to 5 g/kg in mice. No mortality or toxic symptoms were observed during the entire duration of the study [152, 153]. |
| 31    | Citrullus colocynthis        | C. colocynthis plant is safe to use. Studies showed that lethal dose (LD50) to be 200 mg/kg, which indicate that the studied plant is not toxic when comparing the LD50 values of most bioactive pharmaceuticals currently used in therapeutics [154] | In acute toxicity study, there was no mortality observed up to the maximum dose level of 2000 mg/kg body weight of the extract after administered orally [102] |
| 32    | Citrullus lanatus            | Lycopene, vitamin A, cucurbitacin E, citrulline arginine, glutamine and aspartic acid, pectin, vitamin b-complex and minerals [155, 156] | |
| S. No | Names                  | Active ingredients                                                                 | Toxicological evaluation                                                                 |
|-------|------------------------|-------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|
| 33    | Coccinia grandis       | Cephalandrol, β-sitosterol, cephalandrin A and B, β-amyrin acetate, lupeol, cucurbitacin B, taraxerone, taraxerol, β-carotene, lycopene, cryptoxanthin, xyloglucan, carotenoids, β-sitosterol, stigma-7-en-3-one, lupeol, β-amyrin, β-sitosterol, taraxerol [157] | Jatroplphol, jatropha factor C1, C2, C3,C4, C5, C6, jatroplpholones A, B, palmarumycin CP1, JC1, JCV2, curcin, curcacycline A, curcain, β-amyrin, β-sitosterol, stigmasterol, friedelin, taraxasterol, diamide, pyrimidine-2,4-dione, nobiletin, tomentin [103] | The acute toxicity study indicated that treatment of C. grandis is safe up to 2 g/kg tested on animal [158]. Many researchers have confirmed that J. curcas is highly toxic for animal as well as human. All parts of J. curcas are toxic and toxic compound reported from this plant like lectins, curcin, phorbol esters, phytate, protease inhibitors [103] R. communis extracts given by oral route were safe up to a dose of 2,000 mg/kg/BW and did not show any mortality and toxic effects in the behavior of the treated animals [104]. |
| 34    | Jatropha curcas        | Tannins, saponins, flavonoids, anthraquinone glycosides, and flavonoid glycosides [53] | Rutin, quercetin, gallic acid, icin, icin A, kaempferol-3-O-β-rutinoside, quercetin-3-O-glucoside, quercetin-3-O-rutinoside, psoralen, bergapten, coumarin, oleandric acid, α-galactose, and L-rhamnose, 5, 6,8-epiloganic acid, galactose, cinaroside, α-galactoside, cinaroside, myricetin 3-O-glucoside, kaempferol 3-O-glucoside, quercetin 3-O-glucoside, quercetin 7-O-(6-O-rhamnosyl-glucoside), flavones, aglycones, anthraquinone 2-methoxyphenol, 2-pentylpyridine, vitamin E, quinone, Helioxanthin [168]. No literature available. Recommended for natural products, isolation, biological and toxicological evaluation. | Toxicity of 70% methanolic extract of Ficus carica leaves showed LD₅₀ value of brine shrimp assay was 0.158 mg/ml [162] F. sycomorus methanol extract of stem bark is nontoxic up to the dose of 3000 mg/kg [53] S. indicum ethanol extract is safe, up to dose level of 2000 mg/kg in acute toxicity studies in tested animals [165] |
| 35    | Ricinus communis       | β-rutinoside, genistin acid, linolenic acid, α-pinene, α-thujone, stigmasterol, ricinene, β-sitosterol, lupeol, castor oil [159, 160] | Ferulic acid, quercetin-3-O-glucoside, quercetin-3-O-rutinoside, psoralen, bergapten, coumarin, oleandric acid, α-galactose, α-pinene, β-pinene [161] | No data available |
| 36    | Ficus carica           | Tannins, saponins, flavonoids, anthraquinone glycosides, and flavonoid glycosides [53] | Thelignans, sesamolin, sesamin, pinoselin, lariciresinol, α-globulin, β-globulin, triacylglycerols, oleic, linoleic acids, sesamol, γ-tocopherol, 2-furfurylhthiol, 2-phenylethylthiol, 2-methoxyphenol, 2-pentylpyridine, vitamin E, quinone, sesangolin [163, 164] | No literature available. Recommended for pharmacological and toxicological evaluation. |
| 37    | Ficus sycomorus        | Hemicellulose, xyllose, 1-arabinose, d-glucose, d-galactose, and l-rhamnose, 5, 6,8-epiloganic acid, gardoside, plantamajoside [166] | α-pinene, triacylglycerols, oleic, linoleic acids, sesamol, γ-tocopherol, 2-furfurylhthiol, 2-phenylethylthiol, 2-methoxyphenol, 2-pentylpyridine, vitamin E, quinone, sesangolin [163, 164] | 4.5 Gram seed husk one to four times a day soaked in 150ml of warm milk recommend by WHO. Studies confirmed its side effect like bloating, gas, and allergy. No mortality reported [167] |
| 38    | Sesamum indicum        | Protocatechuic acid, catechin, myricitrin, epicatechin-3-O-gallate, avicularin, quercetin, juglanin, kaempferol, myricetin 3-O-(3′-O-galloyl-rhamnopyranoside), cinaroside, liquiritin, rutin [169, 170] | Ficus carica methanol extract is nontoxic up to the dose of 3000 mg/kg [53] S. indicum ethanol extract is safe, up to dose level of 2000 mg/kg in acute toxicity studies in tested animals [165] | No data available |
| 39    | Plantago ovata         | Helioxanthin [168]. No literature available. Recommended for natural products, isolation, biological and toxicological evaluation. | Protocatechuic acid, catechin, myricitrin, epicatechin-3-O-gallate, avicularin, quercetin, juglanin, kaempferol, myricetin 3-O-(3′-O-galloyl-rhamnopyranoside), cinaroside, liquiritin, rutin [169, 170] | No literature available. Recommended for pharmacological and toxicological evaluation. |
| 40    | Polygonum aviculare L. | Jujuboside B1, christinin A, christinin A1 and A2, lotoside II, catechin, epicatechin, kaempferol 3-O-(6-O-rhamnosyl-galactoside), quercetin 7-O-(6-O-rhamnosyl-glucoside), quercetin 3-O-glucoside, kaempferol 3-O-glucoside [171, 172] | Butanol and water extract of Ziziphus spina-christi up to 100 mg/kg and 5 kg/kg, respectively, in animal model produced no functional or structural disturbances in liver and kidney and no hematological changes [173, 174] B. monnieri extract at the dose of 5,000 mg/kg did not cause any side effects. Similarly doses of 30, 60, 300, and 1,500 mg/kg given for 270 days did not produce any toxicity in rats [176]. | No data available |
| 42    | Ziziphus spina-christi  | Bromine, β-sitosterol, betulinic acid, stigmasterol nicotinone, herpespine, bacosides A, bacopasides (I, II, III, IV, V), pseudojujubogenin glycoside, saponins (A, B, C) [175] | Lyciumate, cyclopentapyrrolidine, imidazole, piperidine, nortropane, tropane, pyrrole, spermine, costunolide, cephalmadrins, plantamajoside, quercetin 3-O-glucoside, kaempferol 3-O-glucoside, quercetin 7-O-(6-O-rhamnosyl-glucoside), quercetin 3-O-glucoside, kaempferol 3-O-glucoside [171, 172] | Reported data revealed that LD₅₀ of the L. shawii extract was greater than 2000 mg/kg b.w in animal model [179]. |
| 43    | Bacopa monnieri        | Lyciumate, cyclopentapyrrolidine, imidazole, piperidine, nortropane, tropane, pyrrole, spermine, costunolide, catechin, lyciumaside, emodin, betulinic acid, β-sitosterol glucopyranoside, quercetin, gallic acid, rutin, ρ-coumaric acid, ferulic acid [177, 178] | Solanum nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] | S. nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] |
| 44    | Lycium shawii         | Chlorogenic acid, quercetin, naringenin, solasodine, solamargine, solasoline, α-solanigrine, β-solanigrine, ascorbic acid, nigrummins I and II [180] | Solanum nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] | S. nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] |
| 45    | Solanum nigrum        | Chlorogenic acid, quercetin, naringenin, solasodine, solamargine, solasoline, α-solanigrine, β-solanigrine, ascorbic acid, nigrummins I and II [180] | Solanum nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] | S. nigrum extract at a dose of 2000 mg/kg p.o. was safe and showed no changes/alteration in normal behavior in animal model. No mortality was observed [181] |
Table 2: Continued.

| S. No | Names | Active ingredients | Toxicological evaluation |
|-------|-------|--------------------|--------------------------|
| 46    | Withania somnifera | Withanolides, withaferin, withaferin A, withanone, withanolide A, withanolide IV, withanolide V, withanolide D [182]. | LD_{50} value of W. somnifera extract in rates was greater than 2000 mg/kg body weight. Compared to the control group in subacute toxicity study, administration of extract did not show any toxicologically significant treatment-related changes in clinical observations, and the toxicological studies revealed that the reasonable doses of W. somnifera are nontoxic and safe [182, 183]. |
| 47    | Lantana camara | Eicosane, squalene, β-ionone, Caryophyllene oxide, β-caryophyllene, hexanoic acid, tiglic acid, lantanilic acid, camaric acid, lantadene B, oleanolic acid, lantadene A, lantaninilic acid, lantoic acid, ursolic acid, betulinic acid [184] | L. camara extracts at dose level of 2000 mg/kg and 5000 mg/kg body weight in mice and rats, respectively, showed no significant toxic signs or mortality [185, 186]. |
| 48    | Peganum harmala | Harmine, harmaline, harmalol, harman, vascine and vasicine, pegamine, acacetin 7-O-rhamnoside, 7-O-6″.O-glucosyl-2″-O-(3″-acetylthamnosyl) glucoside, 7-O-(2″-O-rhamnosyl-2‴-O-glucosylglucoside), peganone 1 and 2, p-cymene, limonene, eugenol, thujico acid, β-cubebene [187–189] | P. harmala different doses of different extracts in animal and human clinical studies confirmed that this plant showed side effect like intoxications, abdominal writhing, body tremors, and toxic at high does level causing paralysis, liver degeneration, euphoria, convulsions, nausea, vomiting, hypothermia. However, therapeutic doses have been reported to be safe in a rodent model [189]. |
| 49    | Tribulus terrestris | Naringin, rutin, hyperoside, quercitrin, naringenin, quercetin, hesperetin, kaempferol, apigenin, pyrogallol, gallic acid, catechin, catechol, chlorogenic acid, caffeic acid, vanillic acid, ferulic acid, salicylic acid, ellagic acid, coumaric acid, cinnamic acid [190] | T. terrestris extract showed no mortality/or toxicity at a dose up to 1 g/kg of bw in mice [190]. |
| 50    | Urtica dioica | Anthracin, β-amyrin, β-sitosterol, stigmasterol, oleanolic acid, ursolic acid, quercetin, rutin, chlorogenic, and 2-O-caffeoyl malic acid expressed as caffeic acid, isoquercetin, kaempferol 3-O-rutinoside [191–193]. | U. dioica extracts up to dose level of 2000 mg/kg body weight in animal model showed no mortality or changes/alteration in normal behavior [127]. |

Table 3: Selected antidiabetic phytoconstituents and its targeted metabolic pathway.

| Phytoconstituents | Targeted metabolic pathways | Structures |
|-------------------|-----------------------------|------------|
| Catharantheine   | Free radical; our body has a defense system containing several enzymes, which are catalase, superoxide dismutase, and glutathione-S transferases and reduced glutathione. Catharantheine activates these free radical scavenging enzymes and prevents our body from their adverse effects. | ![Catharantheine](image) |
| Harmine           | Insulin secretion and β-cell regeneration | ![Harmine](image) |
| Betaine           | Carbohydrate digestion and absorption | ![Betaine](image) |
| Achyrantheine β-ecdysone | | ![Achyrantheine β-ecdysone](image) |
| Phytoconstituents                        | Targeted metabolic pathways                                                                 | Structures                                      |
|-----------------------------------------|---------------------------------------------------------------------------------------------|-------------------------------------------------|
| Apigenin                                | Cholesterol synthesis, glycogen synthesis                                                  | ![Apigenin structure](attachment://apigenin.png) |
| Betaine choline                         | Regeneration of pancreatic $\beta$ cells and insulin secretion                            | ![Betaine choline structure](attachment://betaine.png) |
| Ferulic acid                            | Free radical scavenging activity, insulin secretion                                         | ![Ferulic acid structure](attachment://ferulic.png) |
| Leucine, isoleucine, alanine             | Insulin secretion                                                                          | ![Leucine structure](attachment://leucine.png)    |
| Chlorogenic acid                        | Krebs cycle                                                                                | ![Chlorogenic acid structure](attachment://chlorogenic.png) |
| Emodin, cinnamic acid                   | Insulin secretion                                                                          | ![Emodin structure](attachment://emodin.png)     |
| Eugenol, $\alpha$-pinene, limonene, $p$-cymene thujone | Insulin secretion, regeneration of pancreatic $\beta$ cells                              | ![Eugenol structure](attachment://eugenol.png)    |
| Pectin                                  | Glucose transport, carbohydrate metabolism, stabilizing agents                             | ![Pectin structure](attachment://pectin.png)     |
**Table 3: Continued.**

| Phytoconstituents                          | Targeted metabolic pathways                                           | Structures |
|-------------------------------------------|-----------------------------------------------------------------------|------------|
| Inulin                                    | Glucose transport, carbohydrate digestion and absorption              | ![Inulin Structure](image1) |
| Cucurbitacin B                            | Insulin secretion, glycogen synthesis                                | ![Cucurbitacin B Structure](image2) |
| Catechin                                  | Scavenging activity                                                  | ![Catechin Structure](image3) |
| Epicatechin                               | Insulinomematic                                                      | ![Epicatechin Structure](image4) |
| Naringin                                  | Glycogen synthesis, glycolysis, gluconeogenesis                      | ![Naringin Structure](image5) |
| Flavones                                  | Insulin secretion                                                    | ![Flavones Structure](image6) |
| Quercetin, quercitrin, apigenin, rutin,    | Insulin secretion                                                    | ![Quercetin Structure](image7) |
| apigenin-7-O-glucoside                    |                                                                       | ![Quercitin Structure](image8) |
| Naringenin                                | Insulin secretion                                                    | ![Naringenin Structure](image9) |
From the study, the levels of these metals differed in the same plant collected from different geographical locations. A study conducted showed that the levels of lead in Cassia alata varied from 17.7 to 4.45 μg/g for the 5 collection sites. Similarly for Cassia occidentalis and Rauvolfia vomitoria, the level is varied between 7.85-4.35 and 9.25–1.55 μg/g, respectively. Similarly, that of aluminum varied between 105.53 and 23.3 for Rauvolfia vomitoria and 104.25–12.4 μg/g.

| Phytoconstituents                          | Targeted metabolic pathways                                      | Structures                  |
|--------------------------------------------|----------------------------------------------------------------|-----------------------------|
| α-Terpineol                                | Insulin secretion                                               | ![Image](image1)             |
| Kaempferol, isorhamnetin, caffeine acid, p-coumaric acid | Free radical scavenging activity Carbohydrate digestion and absorption, insulin secretion | ![Image](image2)             |
| Vitamin A, E                               | Free radical scavenging activity                               | ![Image](image3)             |
| Ellagic acid                               | Carbohydrate digestion and absorption, insulin secretion        | ![Image](image4)             |
| Carvacrol, linalool                        | Insulin secretion, carbohydrate digestion and absorption        | ![Image](image5)             |
| Stigmasterol                               | Regeneration of pancreatic β cells, insulin secretion           | ![Image](image6)             |
| Ursolic acid                               | Regeneration of pancreatic β cells and insulin secretion        | ![Image](image7)             |
| β-Sitosterol                               | Insulin receptor (IR) and glucose transporter 4                 | ![Image](image8)             |
for Paullinia pinnata. The levels of heavy metals also varied for different plants collected from the same location. Uptake of metals by plants is influenced by a number of factors including metal concentrations in soils, cation exchange capacity, soil pH, organic matter content, types and varieties of plants, and plant age. However, the prevailing factor is the concentration of the metal in the soil and thus the existing environmental conditions [209]. Another study conducted on onion bulb showed that the concentrations of Cr in onion bulb and Fe in onion leaf were above the permissible level (2.3 mg/kg, 425.5 mg/kg) set by FAO/WHO at Mojo (4.87 mg/kg, 1090.40 mg/kg), Meki (4.13 mg/kg, 1836.47 mg/kg), and Ziway (3.33 mg/kg, 764.33 mg/kg), respectively. The results generally indicate that the consumption of these onion bulbs could be the health risk respective to Cr [210]. Therefore, it is suggested that the medicinal plant source for the treatment of diabetes must not be taken from heavy metal contaminated areas to avoid their uptake by the plants because migration of these contaminants into non-contaminated areas (or leaching through the soil and spreading of heavy metal contaminated sewage sludge) are a few examples of events contributing to contamination of the ecosystem.

5. Conclusion and Recommendations

The present review provides a picture of medicinal plants that have been studied as anti-DM drugs, which can be grown either in combination with other medicinal plants or alone as treatment for diabetes and drawbacks should be properly addressed so that medicinal plants can be effectively utilized as anti-DM drugs. Diabetes is a metabolic disorder which can be considered as a major cause of high economic loss which can in turn impede the development of nations. Moreover, uncontrolled diabetes leads to many chronic complications such as blindness, heart failure, and renal failure. In order to prevent this alarming health problem, the development of research into new hypoglycemic and potentially antidiabetic agents is of great interest. In conclusion, this paper has presented a list of anti-DM plants used in the treatment of diabetes mellitus. Bioactive antidiabetic phytoconstituents which showed that these plants have hypoglycemic effects and highly recommended for further pharmacological purposes and to isolate/identify anti-DM active agents also need to investigate the side effects of active ingredients.

Data Availability

This is a review article. All data are taken from published research papers and available online.

Conflicts of Interest

The authors declare no conflicts of interest.

Authors’ Contributions

All three authors contributed equally.

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