Unani formulations for management of diabetes: An overview

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Abstract

Diabetes is a leading cause of death. In India, total number of diabetic patients is expected to be 79.4 million in 2030. It is a multifactorial disease leading to several complications. Traditional medicines are being looked up once again for its treatment; the WHO also recommended its further investigation. In this review, these drug its pharmacological activity and its formulation in Greco-Arab or Unani Medicine in India is explored. Single-drug used in formulation with reported anti-diabetic and related beneficial property are Gymnema sylvestre, Azadirachta indica, Aloe vera, Momordica charantia, Acacia arabica, Eugenia jambolana, Trigonella foenum graecum, Punica granatum, etc. Phytomedicine used in Unani medicine presents an exciting opportunity for the development of new types of therapeutics for diabetes mellitus with a holistic approach, which includes various photochemical groups such as alkaloids, terpenes, and phenolics. Beside hypoglycemic activity, these drugs also have antioxidants, bitter and other activity related to disturbance in carbohydrate metabolism. They are used in the form of decoctions, infusions, tablets, pills, and powder such as Qurse Dhayabitus, Qurse Tabasheer, Safaof Gilo, Qurse Marwareed, Qurse Gulnar, Dawaul Misk Talkh, Sharbate Afseenteen, Roghane Qusht, and Ma-Ussheer and common marketed formulation is Dolabi tablets, Qurs Kushta baiza murgh, Qurs Kushta Zamarrud, Qurs Ziabetus, Safaof Ziabtes, Diab-eaze, etc. There is the tremendous scope of Unani single and compound formulations in the comprehensive management of diabetes particularly Type-2 diabetes; it can also be utilized as an adjuvant with the conventional drug due to its diverse and related beneficial pharmacological activity in diabetes.

Key words: Comprehensive, diabetes, formulations, management, Unani medicine

INTRODUCTION

Diabetes is one of the five leading causes of death.[¹] Diabetes mellitus (DM) is a metabolic disorder complex in nature, resulting in either insulin insufficiency or insulin dysfunction. It is of two types, that is, Type I diabetes (insulin dependent) and Type II diabetes (noninsulin dependent), constituting 90% of the diabetic population.[²] The prevalence of diabetes for all age-groups worldwide was estimated to be 2.8% in 2000 and expected to be 4.4% in 2030. In India, the total number of diabetic patients was 31.7 million in 2000 and is expected to be 79.4 million in 2030. The urban diabetic population in developing countries is projected to double from 2000 to 2030. In another study, Akhter et al. showed that 8% Indian population is diabetic.[³,⁴] Globally, 422 million adults were estimated as diabetic in 2014 and have nearly doubled since 1980, rising from 4.7% to 8.5% in the adult population. Diabetes caused 1.5 million deaths in 2012. Percentage of deaths before the age 70 is higher in low-income and middle-income countries than in high-income countries.[⁵]

Diabetes has become more common in the past few decades due to the stress of growing population, ecological disturbances, and rapid changing lifestyles.[⁶] Type 2 DM (T2DM) has now...
It is currently suspected that diabetes is a multifactorial disease leading to several complications.

Pathophysiology of diabetes is not fully understood, but experimental evidence suggests the involvement of free radicals in the pathogenesis of diabetes and multiple abnormalities of lipoprotein metabolism in the development of diabetic complications. It is currently suspected that hormones produced in fat around abdominal organs are a precipitating cause of insulin resistance which can contribute to a diagnosis of Type 2 diabetes. Diabetes is a multifactorial disease leading to several complications.

Unani medicine literature described diabetes symptomatically as characterized by excessive thirst with increased frequency of urination. It was described by various Greek physicians, and word diabetes is derived from Greek word “Diabanein” which means to “passing through” or “run through” as stated by Ibn Sina in his treatise “Al qanoon fit tib.” The causes of diabetes mentioned are disordered temperament of the weakness of kidney, bladder, and liver, termed as Sue Mizaj wa Zaufe Kulliya, masana wa Jigar.

**TRADITIONAL / UNANI (GRECO-ARAB) MEDICINE FOR DIABETES**

Behind conventional anti-diabetic drugs, antioxidants and multiple therapeutic approach for disturbance in carbohydrate metabolism are under consideration nowadays for treatment of diabetes. Traditional medicine is being looked up once again for the treatment of diabetes, near about 400 traditional plants drug treatments for diabetes have been reported, and on few drug efficacy evaluations for their hypoglycemic activity have been done on animal and human model in Type 2 diabetes. The World Health Organization recommended that traditional medicinal herbs be further investigated. Diabetes leads to various complications such as blindness, kidney failure, coronary artery disease, and gangrene of lower extremities. Due to to these complications, the researchers of a different system of medicine are concentrating on the development of the new anti-diabetic drugs. Anti-diabetic drugs of contemporary medicine have potent and effective hypoglycemic action, but the long-term use of these drugs results in the development of various adverse effects. Therefore, there is a dire need to develop safe and effective drug for the management of Ziabetes Shakri (DM).

**PLANTS WITH ANTI-DIABETIC AND RELATED BENEFICIAL EFFECTS IN INDIAN SYSTEM OF MEDICINE (ISM)**

Plants with reported antidiabetic and related beneficial property after animal and clinical research are: Gymnema sylvestre, Azadirachta indica, Aloe vera, Momordica charantia, Acacia arabica, Aegle marmelos, Allium cepa, Allium sativum, Althaea officinalis, Caesalpinia bonducuella, Cinnamomum zeylanicum, Emblica officinalis, E. jambolana, Ficus racemosa, Plantago ovate, Trigonella foenum graecum, Tinospora cordifolia, Panica granatum, etc. Still, further clinical research is required for the development of the antidiabetic traditional drug. Phytomedicine used in traditional medicine presents an exciting opportunity for the development of new types of therapeutics for DM, which includes various photochemical groups such as alkaloids, terpenes, and phenolics. Several active compounds have been isolated from the plant and herb species of ISM. They are dietary fibers, alkaloids, flavonoids, saponins, amino acids, steroids, peptides, and others. These constituents have produced potent anti-hyperglycemic, hypoglycemic, and glucose suppressive activities. They act by either insulin release from pancreatic β-cells, inhibited glucose absorption in the gut, stimulate glycogenesis in liver or increased glucose utilization by the body. These constituents also exhibited antioxidant, hypolipidemic, and anticitartar activities; they restore enzymatic functions, cause repair and regeneration of pancreatic islets and the alleviation of liver and renal damage. Some active constituents obtained from plants possess insulin-like activity and can be an alternate for insulin therapy.

**UNANI DRUGS / FORMULATIONS USED IN DHAYABITUS (DIABETES)**

Unani system of medicine can be a very rich source as a phytomedicine, many single drugs, and compound formulations are used in Unani medicine for the treatment of Dhayabitus (diabetes). Phytomedicine used in Unani medicine presents an exciting opportunity for the development of new therapeutics / formulations for DM, which include various photochemical groups such as alkaloids, terpenes, and phenolics. Besides hypoglycemic activity, these drugs also have antioxidants, bitter, and other activity related to a disturbance in carbohydrate metabolism. They are used in the form of decoctions, infusions, tablets, pills, powder, confection, etc.

Example of single drugs used for the treatment of Dhayabitus (diabetes) in Unani medicine is Gule surkh, Gulnaar, Roghane gul, Roghane Neelufar, Aabe Jangali Kaasni, Gile Armani, Sandal Safed, Tukhme Khurfa, Tukhme Kahu, Rubb Angoor Khaam, Aabe Khurfa Sabz, Loabe Isappghol, and Kishneez Khushk. Commonly used formulations for diabetes are...

Safoof Ziabetus also have many variants in several texts of Unani medicine, such as it contains Gudmar boti, Soonth, and Jamun.[24] Gudmar booti and Jamun chal (bark); Gudmar booti, Jamun, Afyun, Bisbasa, and Ilaichi kalan.[25] Gilo, Gudmar, and Shaker(sugar)[26]. Its variants are named as Safoof Ziabetus Sada, Safoof Ziabetus Dulabi, and Safoof Ziabetus Qawi.[22]

Qurse tabasheer is commonly used in Unani medicine for diabetes, different category of Qurse tasheer is mentioned in Unani text with different ingredients and indications including diabetes, these are as follows: Qurse Tabasheer, Qurse Tabasheer qabiz, Qurse Tabasheer gulnari, Qurse Tabasheer Afyuni, Qurse Tabasheer Raazi, Qurse Tabasheer kafoori, Qurse Tabasheer Kafoori mulaiyan, Qurse Tabasheer Loolooi sahgee; Qurse Tabasheer mushil, Qurse Tabasheer mulaiyan, Qurse Tabasheer kafoori loolooi, and Qurse tabasheer sartani.[18,21,27,28] Different formulae of Qurse Tabasheer: Several formulations with the different ingredient in the name of Qurse Tabasheer alone are mention in Unani text. Formulation mention in the name of Qurse Tabasheer in Al-Qarabadeen,[18] Qarabadeene Najmul Ghani,[20] Qarabadeene Majeedi,[21] and Kamillus Sana contains different ingredients,[14] whereas formulation of Qurse Tabasheer mention in Bayaaze Kabeer[29] Kitabul Murakkabat Al Marof Makhzan ul Murakkabat,[30] and Kitab Al Murakkabat, [31] contains same ingredients, that is, Tabasheer, Gule Surkh, Gulnar, Tukhme kahu, Tukhme khurfa, and Gile Armani and is indicated in the treatment of Dhayabitus (Diabetes) and its dose mention is 5 g.

COMMON MARKETED FORMULATION OF UNANI FORMULATION FOR TREATMENT OF DIABETES

Dolabi tablets, Qurs Tabasheer, Qurs Kuhsa baize margh, Qurs Kuhsa Zamaraul, Garlitab, Qurs Ziabetes, diabetic care tablets, Gurmar, Neem and Fenugreek capsule, Hoodiab capsule, Jawarish Zarooni Sada, Jawarish Mastagi sada, Shugar no, Safoof Ziabetes dulabi, Herbo diaecon, and Diab-eaze,[32] present marketed formulations are mentioned in Table 1.

Common ingredients present in commonly marketed 23 anti-diabetic formulations of Unani medicine / Pharmacy in decreasing order of their presence in the formulation are Jamun (E. jambolana) - 10, Gudmar booti (G. sylvestre) - 9, Tabasheer - 6, Aqiaia / Babool (A. arabica) - 6, Methi / Halba (T. foenum-graecum) - 5, Karela (M. charantia) - 5, Kusha baiza margh (ash of hen’s egg shells) - 4, Gilo (T. cordifolia) - 4, Guilar (P. granatum) - 4, Kallonji seeds (Nigella sativa) - 3, Shilajeet - 3, Tukhme kahu (Lactuca sativa) - 3, post Gular (F. racemosa) - 3, Chiraita talkh (Swertia chirayita) - 3, Neem (A. indica) - 3, Tukhme khrufa (Portulaca oleracea) - 2, Gule-surkh (Rosa damascena) - 2, Camphor (Kafoor) - 2, Amla (E. officinalis) - 2, Rubb-us-soos (Glycyrrhiza glabra) - 1, Zanabeel (Zingiber officinalis) - 1, etc. Out of surveyed formulations, only three were found to be pharmacopeial formulations rest were patient and propriety formulations.

VALIDATION WORK DONE ON SOME UNANI FORMULATIONS IN DIABETES

Qurse Tabasheer

Effect of Qurs Tabasheer was assessed in streptozotocin (STZ) 60 mg/kg, i.p. single shot, induced diabetic Wistar rats; level of hexokinase and glucose-6-phosphatase was decreased to a significant level while the level of fructose-1-6-biphosphatase was augmented. It significantly reduces the level of serum glucose, total cholesterol, triglycerides, glucose-6-phosphatase, and fructose-1-6-biphosphatase, while the magnitude of high-density lipoprotein cholesterol and hexokinase was amplified anti-hyperglycemic, anti-hyperlipidemic activity of Qurs Tabasheer extract was found to be more effective than standard oral hypoglycemic drug glimepiride.[33] Daily oral administration of Qurse Tabasheer (1 g/kg/day) for 7 days increased glucose tolerance in albino rats. After glucose loading, maximum effect was observed in 90 min, but it was found statistically insignificant.[34]

Triphala Formulation

Terminalia chebula, Terminalia bellerica, and E. officinalis: All three components of Triphala showed significant anti-diabetic properties individually and in combination (rat model of insulin resistance), it inhibited 50% of lipid peroxidation induced with Fe2+/ascorbate was food to be 85.5, 27, 74, and 69 mug/ml, respectively. The concentration needed for the inhibition of hydroxyl radical scavenging was 165, 71, 155.5, and 151 mug/ml, and that for superoxide scavenging activity
<table>
<thead>
<tr>
<th>S. No.</th>
<th>Name</th>
<th>Dosage</th>
<th>Ingredients</th>
<th>P/P and Pharmacy</th>
<th>Manufacturing industry/pharmacy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Safoof-e-ziabetis</td>
<td>3–6 g bid.</td>
<td>Gudmar booti, jamun</td>
<td>P/P</td>
<td>Hermas Unani Herbal Calicut, India</td>
</tr>
<tr>
<td>2.</td>
<td>Cap zиabetis</td>
<td>1–2 cap bid.</td>
<td>Darchini, pan bhadana</td>
<td>P/P</td>
<td>Hermas Unani Herbal Pharmaceuticals, Kerala, India</td>
</tr>
<tr>
<td>3.</td>
<td>Cap ziyabaneel</td>
<td>1–2 cap bid.</td>
<td>Gunde-babool, bark of babool, camphor</td>
<td>P/P</td>
<td>Hermas Unani Herbal Pharmaceuticals Kerala, India</td>
</tr>
<tr>
<td>4.</td>
<td>Shакrino</td>
<td>2 tablets bid.</td>
<td>Aqaqia, tabasheer, shilajit, Gudmar booti, maghze-jamun, post anar, post gular, kushtha khabulsadeed, kushtha marjan, kushtha baize-e-murgh, Kushta sadaf, warq nuqra</td>
<td>P/P</td>
<td>Rex remedies Delhi</td>
</tr>
<tr>
<td>5.</td>
<td>Kalonji sugar powder</td>
<td>1 tsf tid.</td>
<td>Kalonji powder, tukhm-e-jamun, Gudmar booti, tuqme-katayla, tuqm-e-kasni, tuqme-methi</td>
<td>P/P</td>
<td>Mohammedia Products, Karimnagar, India</td>
</tr>
<tr>
<td>6.</td>
<td>Diabeat</td>
<td>1–2 cap bid.</td>
<td>Tukhm-e-kalonji, methi seeds, tukm-e-kasni, neeb</td>
<td>P/P</td>
<td>Hamdard laboratories, Gurugram, India</td>
</tr>
<tr>
<td>7.</td>
<td>Diabetoz</td>
<td>2 cap bid.</td>
<td>Asgand, banslochan, chirahta, gilo, Gudmar, HalelaZard, kolanji, jamun, neem, methi seeds, zanjibeeel</td>
<td>P/P</td>
<td>Hakeem Baqai’s Medicare (p) Ltd.</td>
</tr>
<tr>
<td>8.</td>
<td>Dolabi</td>
<td>1 tab bid.</td>
<td>Gudmar-booti, jamun, kushta-baiza-murgh, tukhm-hummaz, aqaqiya, labbabuz, banslochan, kushtajast, kushtha khubsul-hadeed, gond safaid</td>
<td>P/P</td>
<td>Hamdard laboratories, Gurugram, India</td>
</tr>
<tr>
<td>9.</td>
<td>Gurmar capsules</td>
<td>1 cap bid.</td>
<td>Gudmar booti dried extract</td>
<td>P/P</td>
<td>Dehlvi naturals, Delhi, India</td>
</tr>
<tr>
<td>10.</td>
<td>Jamun Sirka</td>
<td>10–15 ml/day</td>
<td>Jamun fruit pulp, water</td>
<td>P/P</td>
<td>Dehlvi naturals Delhi</td>
</tr>
<tr>
<td>11.</td>
<td>Kerala capsules</td>
<td>2 cap. daily morning</td>
<td>Karela (bitter melon) dried extract, Karela powder</td>
<td>P/P</td>
<td>Dehlvi naturals Delhi</td>
</tr>
<tr>
<td>12.</td>
<td>Karelajamunras</td>
<td>10–15 ml bid.</td>
<td>Karela (bitter melon), Jamun</td>
<td>P/P</td>
<td>Dehlvi naturals Delhi</td>
</tr>
<tr>
<td>13.</td>
<td>Methi capsules</td>
<td>2 cap bid.</td>
<td>Methi dried extract, methi powder</td>
<td>P/P</td>
<td>Dehlvi naturals Delhi</td>
</tr>
<tr>
<td>14.</td>
<td>Qurs-Tabasheer</td>
<td>5 g</td>
<td>Tabasheer, tukhmeh-khurfa, tukhmeh-kahu, gule-surkh, guinar, gile-armani[g39]</td>
<td>Ph</td>
<td>Hamdard laboratories, Gurugram, India</td>
</tr>
<tr>
<td>15.</td>
<td>Kerala Ras</td>
<td>10 ml bid.</td>
<td>Karela (bitter melon)</td>
<td>P/P</td>
<td>Dehlvi naturals Delhi</td>
</tr>
<tr>
<td>16.</td>
<td>Qurs Ziabitis</td>
<td>2 tab bid.</td>
<td>Tukhmeh-khurfa, tukhmeh-kahu, rub-us-soos, tabasheer, gile-armani, gul-e-surkh, kishneezhusk, aqaqia, samag-e-arabi, sandal safaid, sandal surkh, guinar, camphor (NFUM)</td>
<td>Ph</td>
<td>Dawakhana tibbiya college AMU, Aligarh, India</td>
</tr>
</tbody>
</table>

(Contd...)
was found to be 20.5, 40.5, 6.5, and 12.5 μg/ml, respectively. Extracts in 100 mg/kg body weight on oral administration significantly reduce the blood sugar level in 4 h in normal as well as in alloxan (120 mg/kg) induced diabetic rats. Continued, daily administration of the drug produced a sustained effect. Anti-diabetic activity of the formulation and its relationship with their antioxidant property is displayed in the work.\(^{[23]}\)

**Safoof Ziabetus**

*Safoof* (Powder) containing Gurmar Booti (*G. sylvestre*) and Gilo Khushk (*T. cardiofolia*) was studied, showed significant response on blood sugar level in patients average reduction of 56.83 mg/dl (32.80%) in fasting and 67.23 mg/dl (26.94%) in postprandial blood sugar (PPBS) was observed (\(P < 0.05\)).\(^{[35]}\)

**Capsule Gurmar**

It is a combination of powders and aqueous dry extracts of *G. sylvestre, M. charantia, T. foenum-graecum, Coccinia cordifolia,* and *Lagerstroemia speciosa.* Study showed the presence of carbohydrates, glycosides, saponins, steroids, flavonoids, alkaloids, and tannins in the formulation. It exhibited high total antioxidant capacity (622.326 mg/g) and had high flavonoid (386.43 mg/g) and phenol (184.60 mg/g) contents. Hypoglycemia activity of the formulation was comparable to that of the standard oral hypoglycemic drug, metformin hydrochloride at the dose of 100 mg/kg. Results indicate that capsule Gurmar possesses mild analgesic, antioxidant, central nervous system depressant, cytotoxic and hypoglycemic properties.\(^{[36]}\)

**Dolabi**

Prokinetic activity of Unani herbomineral formulation (Dolabi) in STZ induced diabetic rats and its *in vitro* antioxidant activity. Percentage of gastric emptying (GE) and intestinal transit (IT) was significantly (\(P < 0.05\)) decreased in the diabetic rat as compared to normal control groups. In
STZ-induced diabetic rats, Dolabi significantly ($P < 0.05$) accelerated both GE and IT as compared to diabetic control rat’s exhibit in vitro antioxidant activity.\[37\] Effect of a Dolabi (herbomineral formulation) on early diabetic nephropathy demonstrates that Dolabi has an ability to prevent the progression of early diabetic nephropathy. Such protective effect might be due to the presence of flavonoids (catechin, quercetin, and rutin) and triterpene saponins (oleanolic acid and gymnemic acid) present in ingredients of Dolabi which are known to possess potent antioxidant properties.\[38\]

Hypoglycemic potential Dolabi evaluated with pioglitazone in STZ-induced diabetic rats. Groups receiving Dolabi (35.2 mg/kg/day) and pioglitazone showed significant reductions ($P < 0.05$) in fructosamine levels and fasting plasma glucose and oral glucose tolerance test, Sluggish hypoglycemic effects were noted for Dolabi at manufacturer’s recommended doses but at a higher dose; however, good glycemic control was attained, and the results were comparable to pioglitazone.\[39\]

**Damtab**

Polypharmaceutical Unani tablet contains leaf, bark, and seed of *A. indica*, seed of *Gossypium herbaceum*, leaves of *G. sylvestre*, petals of *Rosa damascena*, and seeds of *Syzygium cumini* as the plant constituent. The animal constituent is *Kushta baiza murgh*, that is, ash of eggshells of hen. Clinical study on Damtab 700 mg shows placebo-like effect whereas Damtab 1400 mg possesses hypoglycemic effect.\[40\]

**Jawarish Zar’uni**

Prepared excluding sugar in the form of capsule exhibited a significant effect in reducing microalbuminuria, 24 h urinary protein and improved subjective parameters; fatigue, leg edema in diabetic nephropathy, test drug is safe, effective in the treatment of diabetic nephropathy.\[41\]

**Diabrid**

Qadri et al.\[42\] evaluated antidiabetic activity of Diabrid comprising *G. sylvestre*, *M. charantia*, *E. jambolana*, and *T. foenum graecum*, showed dose-dependent and gradual hypoglycemic activity with no deleterious effect on kidney and liver.\[42\]

*Qurs-e-Ziabetes* obtained from Hamdard Dawakhana is a polyherbal Unani preparation for T2DM. In a clinical evaluation for its anti-diabetic effects was found to have significant therapeutic effects in T2DM with respect to lowering blood sugar levels as well as clinical improvement.\[43\]

Combination of Unani drugs (Several clinical trials are been conducted to evaluate combined effect of two or more drug): Combined effect of *Withania coagulans* Dunal and *T. foenum-graecum* Linn, reveals that it exhibited hypoglycemic activity and significant improvement in symptoms and signs were observed, and significant euglycemia was attained.\[37\] *Arusa* (leaves of *Adhatoda vasic* Nees.) and *Shoneez* (seeds of *N. sativa* Linn.) studied for the hypoglycemic effect of the individual drug found to be effective ($P < 0.05$). Therefore, the study was designed to find out any synergistic effect of the combination in acute and 1 week models in healthy adult alloxan-induced diabetic albino rabbits of either sex with the standard as glibenclamide. The study revealed that the aqueous extract of test combination given orally reduced the blood glucose level. The significant reduction ($P < 0.05$) in blood glucose level started after 3 h and continued for 6 h in both the groups.\[46\] Anas et al.\[47\] thorough review of classical and modern literature selected a formulation containing *Neem, Kalaunji, Karela, Methi, and Jamun* in equal quantity and conducted a clinical trial. The formulation was given in powder form, in 6 g BID doses for 3 months in the diagnosed patient of DM. Formulation exhibited anti-hyperglycemic activity with significant improvement in sign and symptom.\[44\]

Some combinations work more on symptoms of diabetes, for example, clinical study on a Unani formulation consisting of *Satte Gilo* (*T. cardifolia*), Tabasheer (*Bambusa bambos*), and *Maghze Kanwal gatta* (*Nelumbo nucifera*) in management of *Ziabetes shakari* (DM type 2) done by Khan et al. reveals that the test drug exhibited significant effect on subjective parameters such as polyuria and progressive weakness ($P < 0.05$), in polydipsia and unexplained weight loss, and tiredness ($P < 0.001$); while there was no effect in polyphagia ($P > 0.05$). On objective parameters, there was significant effect observed on urine sugar ($P < 0.01$) and glycosylated hemoglobin ($P < 0.05$), while there was no significant effect found on fasting blood sugar and PPBS ($P > 0.05$).\[45\]

Study of diabetic peripheral neuropathy and therapeutic evaluation of Unani formulation *Habee Azaraqi* in its management exhibited significant improvement in pain ($P < 0.05$), numbness ($P < 0.001$), burning sensation ($P < 0.001$), and paresthesia ($P < 0.05$) in subjective parameters in intragroup comparison.\[46\] Itrifal Kishneezi showed considerable in vitro antioxidant activity in a dose-dependent manner commonly used Unani drugs and drug used placed under Bitters / *Musaffie Dam* (Blood Purifiers) category for treatment of *Dhayabitus* (diabetes) in Unani formulations with reported anti-diabetic and other related activity is depicted in Tables 2 and 3.

**DISCUSSION**

Active principles useful in diabetes are present in Unani formulations such as dietary fibers, alkaloids, flavonoids, saponins, amino acids, steroids, peptides, and others. These drug produces potent hypoglycemic, anti-hyperglycemic, and glucose suppressive activities. Effects from the Unani formulations / compounds may be achieved by either insulin release from pancreatic β-cells, inhibited glucose absorption...
### Table 2: Reported anti-diabetic and related activity in drugs used in Unani formulations

<table>
<thead>
<tr>
<th>Unani name</th>
<th>Botanical name</th>
<th>Antidiabetic and other favorable activities/effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tukhme methi</td>
<td><em>Trigonella foenum graecum</em></td>
<td>Insulin stimulating properties/hypoglycemic effect (in diabetic rats, dogs, mice, and healthy volunteers)/(in IDDM and NIDDM clinical trial), significant hypolipidemic, hypocholesterolemic, anti-oxidant activity. Antioxidant activity. 4-Hydroxyloleucine (amino acid) mainly distributed in fenugreek increased glucose-stimulated insulin release by isolated islet cells in rats, mice, and humans.</td>
</tr>
<tr>
<td>Babool (Aqaaqia/Samaghe Arabi)</td>
<td><em>Acacia arabica</em></td>
<td>Plant extract: Secretagogue to release insulin, induces hypoglycemia (in control rats) (powdered seeds), induced-hypoglycemia (normal rabbits) initiating release of insulin from pancreatic beta cells.</td>
</tr>
<tr>
<td>Jamun</td>
<td><em>Eugenia jambolana</em></td>
<td>Kernels antihyperglycemic, shows a reduction in blood glucose level. (pulp) Hypoglycemic activity STZ induced diabetic mice, (extracts) inhibited insulinase activity from liver and kidney. Hypoglycemic effects (Mukherjee et al., 2006), formulation Madeglycyl, prepared from the seeds of <em>E. jambolana</em>, showed alleviation in the symptoms of diabetes in clinical trials (Denis et al., 2008), anti-diabetic features in type 2 diabetes (glycemia rates) return to normal levels within 3–6 months in 75% of cases particularly in obese patients. Glycemia rates start to decline after 15 days; it acts by improving the ability of the body's tissues to absorb glucose, thus enhancing the effectiveness of insulin.</td>
</tr>
<tr>
<td>Anar/Gulnar</td>
<td><em>Punica granatum</em> (Gulnar/Flower)</td>
<td>Flower extracts antioxidant, anti-hyperglycemic effect with reduction in glycosylated hemoglobin levels (STZ-induced DM. in rats) Pomegranate flower has dual PPAR-α/γ activator properties which is a major regulator of lipid and glucose metabolism and is a natural PPAR α and γ agonist. Inhibition of carbohydrate digestive enzymes (α-amylase and a-glucosidase) and their phenolic content may contribute to the anti-hyperglycemic effects of pomegranate flower and peel support its claim in diabetes in Unani and Chinese Medicine.</td>
</tr>
<tr>
<td>Bel</td>
<td><em>Aegle marmelos</em></td>
<td>Leaves reduce blood sugar and urea, serum cholesterol (alloxanized rats). With hypoglycemic activity, its extract also prevented a peak rise in blood sugar at 1h in oral glucose tolerance test.</td>
</tr>
<tr>
<td>Tukhme Kahu</td>
<td><em>Lactuca sativa</em></td>
<td>Hypoglycemic effect, Lactucin and lactucopicrin, isolated from <em>Lactuca scariola</em> have shown hypoglycemic effect (Jaffery and Harborno). Methanolic leaf extract investigated for <em>in vitro</em> inhibition of oxidative damage induced by UV-radiations to the <em>Salmonella typhi</em> bacteria and <em>in vivo</em> effect on the production of body enzymes, that is, catalase and superoxide dismutase. Shown significant antioxidant potential. Antioxidant activity was displayed by ethanolic extracts by means of spectrophotometric methods. Antioxidant potential both <em>in vitro</em> and <em>in vivo</em>, capable of protecting neurons against glucose/serum deprivation-induced cell injury (neuroprotection) have potential in neurodegenerative disorders.</td>
</tr>
</tbody>
</table>

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### Table 2: (Continued)

<table>
<thead>
<tr>
<th>Unani name</th>
<th>Botanical name</th>
<th>Antidiabetic and other favorable activities/effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tukhme Khurfa</td>
<td>Portulca oleracea</td>
<td>Significant hypoglycemic activity effect was observed when entire plant in dried form was administered intragastrically to rabbits at dose of 1.5 and 2.0 g/kg after 8 and 12 h, respectively. Seeds, in a mixture with 7 other plants, administered orally to male rats at a dose of 4.0 g/animal, were active. In vitro antioxidant activity of the methanolic extract of <em>P. oleracea</em> investigated by Sanja et al. by DPPH displayed free radical scavenging activity, reducing power by FeCl₃, nitric oxide free radical scavenging activity, and superoxide scavenging activity by alkaline DMSO method. Hypolipidemic activity (rich source of omega, 3 fatty acids important in preventing heart attack, and strengthening the immune system), hepatoprotective activity, neuroprotective effect and ameliorates diabetic nephropathy through suppression of renal fibrosis and inflammation (diabetic mice).[68,69,70]</td>
</tr>
<tr>
<td>Amla</td>
<td>Emblica officinalis</td>
<td>Decreases lipid peroxidation, antioxidant, and hypoglycemic[59]</td>
</tr>
<tr>
<td>Palas</td>
<td>Butea monosperma</td>
<td>Anti hyperglycemic[53]</td>
</tr>
<tr>
<td>Post Halela</td>
<td>Terminalia chebula</td>
<td>Antibacterial, hypoglycemic[53,71] aqueous extract (fruits) evaluated for its antidiabetic activity in STZ-induced mild diabetic rats and compared with tolbutamide. Oral administration of effective (dose 200 mg/kg body weight) of aqueous extract of <em>T. chebula</em> daily once for 2 months reduced the elevated blood glucose by 43.2% and significantly reduced the increase in HbA1c. Same dose also improves elevated blood lipids as well as decreased serum insulin levels in contrast to the untreated diabetic animals[72] Chloroform extract of <em>Terminalia chebula</em> seed powder in STZ-induced diabetic rats produced dose-dependent reduction in blood glucose comparable with glibenclamide in short-term study. Significant renoprotective activity is observed in <em>T. chebula</em> treated rats. The result indicates a prolonged action in the reduction of blood glucose by <em>T. chebula</em>; this reduction is probably mediated through enhanced insulin secretion from the β-cells of Langerhans or by extra pancreatic mechanism.[73]</td>
</tr>
<tr>
<td>Balela</td>
<td>Terminalia belerica</td>
<td>Antibacterial, hypoglycaemic[53] dried 75% methanolic extract <em>Terminalia belerica</em> fruit in water studied in alloxan induced hyperglycemia and antioxidant defense mechanism in rats, drug prevented hyperglycemia significantly from the 6th day of administration and 54% reduction was observed on the 12th day. Oxidative stress was also significantly lowered by the administration of extract.[74]</td>
</tr>
<tr>
<td>Asgandh</td>
<td>Withania somnifera</td>
<td>Hypoglycemic, hypocholesterolemic[63]</td>
</tr>
<tr>
<td>Aam (Seed)</td>
<td>Mangifera indica</td>
<td>Hypoglycemic activity (may be due to an intestinal reduction of the absorption of glucose).[53] Alcoholic extract of <em>Mangifera indica</em> leaves and kernel seeds is having significant anti-diabetic effect in alloxone-induced diabetes in Wistar rats by stimulating insulin production in the pancreas.[75]</td>
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<tr>
<td>Pyaz</td>
<td><em>Allium cepa</em></td>
<td>Dried onion powder: Anti-hyperglycemic activity in diabetic rabbits, antioxidant and hypolipidemic activity, normalized the activities of liver (diabetic patient), onion juice significantly controlled post-prandial glucose levels.[53]</td>
</tr>
<tr>
<td>Gule Surkh</td>
<td><em>Rosa damascena</em></td>
<td><em>R. damascene</em> methanol extract of the plant on oral administration significantly decreased blood glucose after maltose loading in normal and diabetic rats in a dose-dependent manner. In addition, it inhibited postprandial hyperglycemia comparable to acarbose. <em>R. damascene</em> was found to be a potent inhibitor of the $\alpha$-glucosidase enzyme, anti-diabetic effect may be mediated by inhibition of $\alpha$-glucosidase that suppressed carbohydrate absorption from the small intestine and can reduce the postprandial glucose level.[76]</td>
</tr>
<tr>
<td>Rabbus soos</td>
<td><em>Glycyrrhiza glabra</em> Linn</td>
<td>Flavonoids-enriched fraction prepared from the EtOH extract of <em>Glycyrrhiza glabra</em> L. roots, exhibited PPAR-c ligand-binding activity, was also effective in preventing and/or ameliorating diabetes, abdominal obesity, and body weight gain in KK-Ay mice and/or high-fat-diet-induced obese C57BL/6J mice.[77]</td>
</tr>
<tr>
<td>Kisneez</td>
<td><em>Coriandrum sativum</em> Linn</td>
<td>Gray and Flatt (1999) have found the antidiabetic potential of coriander seeds in STZ-induced diabetic mice. (Gray and Flatt 1997) also observed an increased glucose transport and incorporation into muscle glycogen and six-fold increased insulin secretion in pancreatic b-cells potentiated by coriander seed powder in <em>in vitro</em> studies.[78-80]</td>
</tr>
<tr>
<td>Zanjabeel</td>
<td><em>Zingiber officinalis</em></td>
<td>Sanjay <em>et al.</em> have studied the antidiabetic effect of the juice of <em>Z. officinale</em> (4 mL/kg), p.o. daily) for 6 weeks on STZ-induced type I diabetic rats with particular reference to the involvement of serotonin (5-hydroxytryptamine; 5-HT) receptors in glycaemic control. <em>Z. officinale</em> produced a significant decrease in fasting glucose level and increase in insulin level in diabetic rats. Treatment also caused a decrease in serum cholesterol, serum triglyceride, and blood pressure in diabetic rats. Antidiabetic activity in type I diabetic rats was possibly by involving 5-HT receptors.[81] Ethyl acetate extract of ginger was evaluated for its antioxidant activity in terms of DPPH radical scavenging potential with an IC50 value of 4.59 $\mu$g/ml. Antidiabetic activity was also evaluated by estimating antiglycation potential (IC$_{50}$ 290.84 $\mu$g/ml). Effect of extract to enhance glucose uptake in cell lines were evaluated in L6 mouse myoblast and myotubes. Antibody-based studies in treated cells revealed the effect of EAG in expressing Glut 4 in cell surface membrane compared to control. Activity is initiated by antioxidant, antiglycation, and potential to express or transport Glut4 receptors from internal vesicles.[82]</td>
</tr>
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</table>
Gular (Bark and fruit)  \textit{Ficus racemosa} Linn

Ethanolic extract is anti-diabetic in the experimental animal.\cite{83}  \textit{F. racemosa} bark powder and aqueous extract in STZ-induced diabetic rats, displayed a significant reduction in blood glucose, serum cholesterol, and triglyceride levels. Aqueous extract was more effective.  \textit{F. racemosa} bark has a significant hypolipidemic and hepatoprotective effect besides being a potent antihyperglycemic agent.\cite{84} The fruit, bark, latex, seeds, or leaves of \textit{F. racemosa} plant have been reported to decrease blood glucose levels in diabetic animals.\cite{85}


### Table 2: (Continued)

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<td>Gulo</td>
<td>\textit{Tinospora cordifolia} (Willd) Miers. Roots/stem</td>
<td>Hypoglycemic activity and hypolipidemic activity (alloxan diabetic rats) decreases the blood glucose level and increases glucose tolerance (rodents).\cite{53} Hexane, ethyl acetate and methanol extract of stem on oral administration were found to have potent antidiabetic activity that reduces blood sugar level in STZ-induced diabetic rats. Supplementation of methanol extract significantly decreases the glycosylated hemoglobin level as compare to diabetic control, the insulin and C-peptide levels were improved which shows the regeneration of ( \beta )-cell which secretes insulin, histopathological studies of pancreas of TCS methanol extract treated groups confirm the regenerating capacity of extract.\cite{95}</td>
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<td>Ailwa/Ghekwar</td>
<td>\textit{Aloe vera} and \textit{Aloe barbadensis}</td>
<td>(Extracts increases glucose tolerance in both normal and diabetic rats, hypoglycemic (alloxanized diabetic rats). (Bitter principle) showed hypoglycemic effect in diabetic rats through stimulation of synthesis and/or release of insulin from pancreatic beta cells, anti inflammatory activity (improves wound healing in diabetic mice).\cite{53}</td>
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### Table 3: Bitters drugs with anti-diabetic and related reported actions used in Unani formulations

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<tr>
<td>Gurmar Booti</td>
<td>\textit{Gymnemama sylvestre} (leaves)</td>
<td>Hypoglycemic activity, inhibit glucose uptake in the intestine. Stimulate insulin secretion from mouse cells and isolated human islets \textit{in vitro}, lowered fasting blood glucose, release insulin probably by causing regeneration of pancreatic ( \beta )-cells both \textit{in vivo} and \textit{in vitro}. Gymnemic acids 1-4, guarmarin shows antidiabetic activity.\cite{54,86,87}</td>
</tr>
<tr>
<td>Kalonji</td>
<td>\textit{Nigella sativa}</td>
<td>Seed extracts enhance glucose-induced insulin release from rat-isolated Langerhans islets and basic sub-fraction largely contributes to this stimulatory effect.\cite{88} Improve the dyslipidemia associated with type 2 diabetic patients, exerts lipid-lowering and insulin-sensitizing actions in the rat, immunomodulatory activity.\cite{89,90} \textit{In vivo} treatment with \textit{N. sativa} seed ethanol extract exerts an insulin-sensitizing action by enhancing ACC phosphorylation (major component of the insulin-independent AMP-kinase signaling pathway) and by enhancing muscle Glut4 protein expression.\cite{91} Decoction (aqueous suspension) at a dose 1–2 g/kg/day displayed hypoglycemic, anti-obesity and hypolipidemic action in \textit{Meriones shawii} (desert gerbil).\cite{92} Seed oil significantly reduce blood suger and increase insulin level.\cite{54} Effect of NSO on blood glucose concentrations was studied in STZ diabetic rats and effect of NSO, nigellone, and thymoquinone were studied on insulin secretion of isolated rat pancreatic islets in the presence of 3, 5.6 or 11.1 mM glucose. NSO significantly reduce blood glucose concentrations after 2, 4 and 6 weeks. The blood lowering effect was not paralleled by stimulation of insulin release in the presence of NSO, nigellone or thymoquinone. indicate that the hypoglycemic effect may be mediated by extra-pancreatic actions rather than by stimulated insulin release.\cite{93}</td>
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<tr>
<td>Gulo</td>
<td>\textit{Tinospora cordifolia} (Willd) Miers. Roots/stem</td>
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<td>Tukhme Karela</td>
<td>Momordica charantia Seeds</td>
<td>Potent hypoglycaemic, significant improvement in glucose tolerance in diabetic subjects without any increase in serum insulin levels, increases the renewal of partial cells in the pancreas or may permit the recovery of partially destroyed cells and stimulates pancreatic insulin secretion.[99]</td>
</tr>
<tr>
<td>Neem</td>
<td>Azadirachta indica</td>
<td>Antihyperglycemic activity (STZ treated rats) because of increase in glucose uptake and glycogen deposition in isolated rat hemidiaphragm, anti-bacterial, antimalarial, antifertility, hepatoprotective and antioxidant effects.[98] Aqueous extract of tender leaves reported to be effective in reducing blood glucose, this effect was due to peripheral utilization of glucose and blocking the action of epinephrine on glycogenolysis.[98]</td>
</tr>
<tr>
<td>Chiraita</td>
<td>Swertia chirayita</td>
<td>Stimulates insulin release from islets.[93] Methanolic leaf extracts contain significant antibacterial and anti-diabetic potential, methanolic root has potential antioxidant activity.[97]</td>
</tr>
<tr>
<td>Afsanteen</td>
<td>Artemisia absinthium</td>
<td>Ethanol extract of Artemisia absinthium in alloxan-induced diabetic rats was given in a dose 250, 500, and 1000 mg/kg body weight in distilled water i/p, for 10 days. All elevated blood serum markers were reduced to significant levels at both medium and high doses and also after glibenclamide treatment.[98] Fifteen patients with diabetes mellitus in a preliminary study were treated with one of the species of Artemisia (Artemisia herba-alba Asso. Extract). Results displayed considerable lowering of elevated blood sugar with no adverse effects.[98]</td>
</tr>
</tbody>
</table>

Botanical/Unani name mentioned are as correlated by NFUM[20] and UPI.[100] NFUM: National formulary of Unani medicine, UPI: Unani pharmacopoeia of India, N. sativa: Nigella sativa, NSO: Nigella sativa oil, N. sativa: Nigella sativa

There is tremendous scope in these Unani formulations in the comprehensive management of diabetes as these compounds also exhibited antioxidant and hypolipidemic activity, restored enzymatic functions, repair, and regeneration of pancreatic islets and the alleviation of liver and renal damage as evident by the review. Deficiency of insulin causes disorder of metabolism of carbohydrate, protein and fat, and these drugs are probably used for correction of a defect of liver and stomach as indicated in classical Unani text and improvement of the functioning of liver also improves the metabolism and ultimately can improve the disease condition in DM. Several bitter drugs are used in the Unani formulation for the treatment of DM, dafe Ziabetus (anti-diabetic) activity is found as a direct indication in some of the bitter drug, or it can also be used in accordance to the etiopathogenesis of diabetes and its complications, several of these bitter drugs are termed as Musaffi-i-Dam. Musaffi-i-Dam (Blood Purifiers) drugs make certain action on blood due to which harmful constituents are altered in such constituents, which are then eliminated easily. Besides, this many drugs / formulation are cardio-protective, neuroprotective, and nephroprotective. Unani polyherbal formulations contain several phytoconstituent which can be responsible for other related activity to control diabetic complications.

Etiologic factor implicated in the development and complication of diabetes is the damage induced by free radicals, and hence an anti-diabetic formulation compound with antioxidant properties would be more beneficial, and ingredients of several Unani formulations possess significant antioxidant property. Numerous physicians of the modern system of medicine are prescribing natural compounds containing flavonoids for their antioxidant potential. Furthermore, oxidative stress has a great role in damaging the insulin-producing cells of the pancreas and diabetic complications in DM.[100,101] Probable mechanism and benefit of medicinal plants present in these formulations are from their antioxidant activities. Most of the medicinal plants with anti-diabetic property possess significant antioxidant activity. Vegetables and fruits antioxidant in comparison to synthetic antioxidants can be more effective due to less alternative dispute resolution and are able to decrease the risk of DM. This oxidative stress usually causes or exacerbates chronic hard curable diseases including diabetes.[103]

Many drugs in the formulation such as Jamun act by improving the ability of the body’s tissues to absorb glucose, thus enhancing the effectiveness of insulin. Gudmar and Gilo which is commonly used in marked Unani formulations can cause regeneration of β-cell, many drug causes increased in glucose transport and its incorporation into muscle glycogen and increased insulin secretion or stimulation of insulin secretion. Many drugs also act by suppressing carbohydrate absorption from the small intestine and Unani formulations containing these drugs can reduce the postprandial glucose level for better management of blood sugar. Gulnar is indicated in Unani medicine for the treatment of DM since centuries and mentioned by eminent Unani scholars such as Jurjani[16] and Majoosi[104] have now gain renewed interest.
due to Pomegranate flower which has dual peroxisome proliferator-activated receptor (PPAR)-α/-γ activator properties (which are a major regulators of lipid and glucose metabolism) and are a natural PPAR α and γ agonist. Some drugs in the formulations display extra-pancreatic actions. We can conclude that these formulations are holistic in their approach.

In Unani medicine different formulation is available for treatment / control of diabetes and its complication with a holistic approach and with higher safety margin which is required for the management of this chronic metabolic disorder. These formulations can play an important role in promoting healthy life in diabetics. Constrain related to these formulations is that which this review clearly indicates is that several authentic pharmacopeial preparations are still not marketed and which can be very beneficial if it reaches through as a pharmacopeial preparations. Several works displayed higher dosage for their proper activity in DM, and there is a need for the reconsideration of dosage of this formulation because under-dosing is quite common when marketed formulations are concerned, higher dose limit can be indicated by review of authentic text in this regard. Adjuvant studies on these formulations with contemporary drug or treatment procedure of DM is lacking. These formulations can also be utilized as an adjuvant with the conventional drug due to its diverse and related beneficial pharmacological activity in diabetes and its complications.

CONCLUSION

The present review reveals that there is the tremendous scope of Unani single drugs and compound formulations in the comprehensive management of diabetes (particularly type-2 diabetes), so it can also be utilized as an adjuvant with the conventional drug due to its diverse and related beneficial pharmacological activity in diabetes and on its complications.

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