INTRODUCTION

Management of diabetes mellitus

The aim of therapy in the treatment of diabetes is to control elevated levels of glucose levels, symptoms related to hyperglycemia, and chronic complications. Management of diabetes has been broadly classified into three domains viz., glycemic control, treat associated conditions, and diabetic complications [1]. Diabetes mellitus is a chronic condition with higher levels of glucose in the body. This alteration in glucose homeostasis is due to the dysfunction of insulin hormone, which helps in mobilization of glucose into cells and produce energy. Diabetes mellitus is further classified into two types i.e. type 1 diabetes mellitus (T1DM) and type 2 diabetes mellitus (T2DM) [2]. The glycated haemoglobin (HbA1c) is considered to be the most precise and best standard for judgment of patient diabetes records in recent years because it provides a sign of a 2 to 3-month glycaemia record [3]. Formulated insulin is obtained from different species of animals with little variation from human insulin e.g., porcine and beef insulin has been used in the treatment of severe diabetic conditions [4]. Porcine insulin at the C-terminal amino acid of chain B has alanine and human insulin contain threonine. Beef insulin differs with three different substitutions viz., alanine (B10), valine (A10), and alanine (A8) [5, 6]. These isolated forms of insulin were used when DNA recombinant technology is not available. They were usually avoided due to adverse immunological responses like type I allergy and injection-site lipatrophy [7, 8]. American diabetes association has laid down four criteria for the diagnosis of diabetes.

1. Fasting glucose level ≥ 126 mg/dl (impaired glucose tolerance) and non-fasting plasma glucose ≥ 200 mg/dl
2. Oral glucose tolerance test (OGTT) ≥ 200 mg/dl
3. Hyperglycaemic symptoms (Polyphagia, polyuria, polydipsia)
4. Glycated haemoglobin (HbA1c) ≥ 6.5%

The Management of diabetes mellitus is followed by insulin therapy, oral hypoglycaemic drugs, other chemical agents, and non-therapeutic methods [9]. The present review includes particulars of currently used alopatic medicaments in the treatment of diabetes mellitus worldwide. The main objective to compile the present review systematically to learn and narrate with classifications, mechanisms of actions, efficacy, and common side effects of FDA approved drugs used in antidiabetic therapies as monotherapy, addition therapy, or in combinations. This review also prominence on physiological reactions like effects on insulin resistance and insulin sensitivity, weight gain or loss, developmental risk, or decline in risk for diabetic complications produce by antidiabetic drugs. It also vaguely covers the effects of hypoglycaemic drugs on metabolic activities and elevated biochemical levels, which are commonly reported in diabetes mellitus patients. The major databases like Google Scholar, MEDLINE, Scopus-Elsevier, Directory of Open Access Journal (DOAJ), and Open J-Gate are employed to compile the current review in January 2020. The complete indexation of all the antidiabetic drugs, hypoglycaemic drugs, T1DM and T2DM drugs, oral hypoglycaemic medicaments, types of insulin, and mechanism of action of antidiabetic drugs are brought into play for literature compilation.

Insulin therapy

Insulin therapy is the main course of treatment for T1DM and uncontrolled hyperglycaemia T2DM [1]. The various insulin delivery systems are available for the administration of insulin e.g., portable pen injectors, inhaled insulin, insulin degludec, insulin syringes, and infusion pumps. Insulin is also administered intravenously, intramuscular, and traditionally insulin is administered by a subcutaneous route [10]. Insulin therapy is given to control abrupt glucose changes in the diabetic state. Thus, insulin administered must be similar in pharmacokinetic profile as that of endogenous insulin released from the pancreas, i.e., a rapid rise in blood concentration, short duration of action and rapid clearance [11]. Commercial available insulin preparations are different from human insulin in the amino acid sequences, solubility, and onset/duration of action. Four types of injectable insulin preparations are available with the different onset of actions and prepared with recombinant DNA technology viz., rapid-acting, short-acting, intermediate-acting, and long-acting [12]. Nearly, all the long-acting insulin reduces the levels of glycated haemoglobin (HbA1c) levels to normal limits [13]. Type 1 diabetic patients are treated with insulin therapy after confirmation of disease symptoms. Patients counseling is necessary before prescribing insulin therapy, e.g., amount of food intake in terms of carbohydrate, need for physical activity, and hypoglycaemia effects [14]. Two types of insulin therapy regimens are followed in
the management of diabetes i.e. intensive and conventional therapy. Intensive insulin therapy is also known as flexible insulin therapy, in which current glucose levels and amount of carbohydrate intake in the meal is considered before the administration of human regular insulin [15]. Hypoglycaemia is the most common reported side effect in intensive insulin therapy [16]. In conventional insulin therapy, premixed human insulin (30% regular and 70% NPH) is administered before breakfast and dinner, which is followed by fixed diet plans [17]. The intensive insulin therapy consisted of daily three or more insulin injections and conventional insulin therapy is designed to control glucose levels with one or two insulin injections daily [18]. Intensive insulin therapy does not have diabetic complications related to T1DM, e.g. nephropathy, neuropathy, retinopathy, and microvascular complications when compared with conventional insulin therapy [19]. Diabetic patients, along with complications like coronary artery surgery, acute stroke, stress, and acute myocardial infarction, have a 20% higher risk of death. Intensive insulin therapy in diabetic and non-diabetic patients with critical illnesses or the surgical intensive care unit reduces morbidity and mortality [20]. In the case of childhood diabetes especially in T1DM, intensive insulin therapy is found to be safer [21]. T2DM patients which undergo gastoectomy for cancer treatment, intensive insulin therapy, reduces short term morbidity [22]. In acute neurological injury, intensive insulin therapy shows better control of glucose levels and reduces morbidity [23]. The therapy causes severe hypoglycaemia if regular monitoring of plasma glucose levels is not recorded. Thus, new automated techniques like islet transplantation, insulin delivery, and preservation of endogenous insulin have been developed in recent years [24].

Rapid-acting insulin
A rapid-acting insulin is the safe, very fast onset of action, rapid clearance, and effective for a short duration. Commercially, three injectable forms are available viz., insulin lispro, insulin aspart, and insulin glulisine. A rapid-acting analog does not cause hypoglycaemia or nocturnal hypoglycaemia and shows better control of postprandial glycaemia [25]. These insulin analogs can be administered before or immediately after meals [26]. Insulin lispro is first marketed insulin preparation in the USA. Insulin lispro has alteration in the amino chain-B of insulin, i.e., proline (B-29) and lysine (B-28). This alteration increases the dissolution rate of insulin because the dimeric form of insulin is converted into monomer and rapidly absorbed, but this modification does not change receptor binding affinity [27]. Insulin aspart is another commercially available rapid-acting insulin preparation for human use and made by the substitution of proline (B-29) with aspartic acid. This exchange alters the interaction between monomer-monomer, i.e., aspartic acid (B-28) and glycine (B-23), which cause a decrease in insulin self-aggregation [12, 28]. Insulin glulisine causes quick disposal of endogenous glucose after post-meal and physiological action similar to regular human insulin [29]. It is produced by substitution of lysine in place of asparagine at B-3 and glutamic acid at B-29 replacing lysine [12]. Insulin glulisine in the solution form exists as a monomer without zinc and has rapid absorption. Thus, it is less immunogenic when compared with other rapid-acting insulin but may cause rare insulin autoimmunity syndrome [30]. It is proving to be the most effective and safe insulin preparation for children suffering from T1DM [31].

Short-acting insulin
Short-acting insulin analogs did not show any hazard of severe hypoglycaemia and shown better management of HbA1c, postprandial glucose, and diabetic ketoacidosis [32]. Regular insulin is short-acting insulin made by DNA recombinant technology. It exists as a hexameric form, which is bulky and creates depot upon subcutaneous injection and result delay in transportation through blood [1]. The onset of action of regular insulin is appearing within 30 min and the duration of action is about 2 to 3 h [12].

Intermediate-acting insulin
Intermediate-acting insulin has delay onset of action and absorption. Three types of intermediate-acting insulin have been used clinically viz., NPH (Neutral Protamine Hagedorn or isophane), insulin glargine, and insulin detemir. NPH at neutral pH is a combined product of insulin and stoichiometric prostamine in the isophane ratio, i.e., neither insulin nor protamine found in large concentrations [33]. NPH is poorly soluble insulin product in blood and due to this property, it resulted in delayed onset of action (onset of action is 2-5 h and duration of action is 4-5 h). NPH insulin shows little improvement in HbA1c level when compared with other intermediate-acting insulins [12]. Insulin glargine, intermediate-acting insulin, is having two modifications, i.e., the addition of two arginine molecules at C-terminus of B-chain of human insulin and replacement of asparagine by glycine. This addition of arginine causes a slight increase in the solubility of insulin glargine towards acidic pH and a shift in isoelectric point from pH 5.4 to 6.7 [34].

Insulin glargine has shown a greater affinity towards insulin-like growth factor-1 receptors compared to regular insulin [35]. The onset of action of insulin glargine is slow, with significant activity maintained for 11-24 h [36]. It can be administered any time during a day and significantly reduces HbA1c levels and prevent hypoglycaemic shocks during nights [37]. Insulin detemir is a new long-acting analog that is created by the substitution of myristic acid at B-29 position and removal of threonine from the B-30 position [38]. These modifications cause self-aggregation and protein binding with albumin. The duration of action of insulin detemir is more than 12 h and having 1-2 h onset of action [36]. Insulin detemir is considered to be much safer than NPH [12]. It shows better control on glycaemia in T1DM because it does not precipitate after subcutaneous administration, binds to albumin protein in the blood, and forms depot, which can be buffered easily if a change occurs in absorption parameters [39].

Mixtures of insulin
A mixture of insulin is a formulation created by mixing rapid and short-acting insulin in the same syringe before administration such as insulin lispro, aspart, glulisine with NPH. Other mixtures of insulin are NPL (Neutral protamine Lispro) and NPA (Neutral Protagmine Aspart) [12]. Insulin preparation containing 50% of short-acting insulin and 50% of regular insulin. These mixtures are well-tolerated, provide efficient control over glycaemia, and can be administered once or twice daily [40]. The premixed formulation, like 70% NPH and 30% regular insulin, improves efficacy, accuracy and gives better control over glycaemia [41].

Oral antidiabetic agents
The oral antidiabetic drugs are used for the treatment of T2DM, which is not controlled with diet restriction and physical exercise. Six categories of oral antidiabetic agents have been available in the market viz., biguanides, sulfonylureas, meglitinides, thiazolidinediones, alpha-glucosidase inhibitors, dipeptidyl peptidase-IV (DPP-4) inhibitors, sodium-glucose co-transporter-2 (SGLT2) inhibitors, other agents and non-therapeutic methods [12, 62]. The first-line therapy for T2DM treatment is started with the most widely used oral antidiabetic drug i.e., metformin. The second line of therapy involves sulfonylureas, thiazolidinediones, glucagon-like polypeptide-1 (GLP-1) agonists, dipeptidyl peptidase-IV (DPP-4) inhibitors, meglitinides, and SGLT2 inhibitors [43].

Biguanides
Metformin, a dimethyl biguanide, is used in the treatment of T2DM chronically without an increase in the risk of hypoglycaemia [44]. Its efficacy, tolerance, safety, improved peripheral insulin sensitivity without an increase in insulin secretion, weight gain, and other beneficial effects on cardiovascular systems promote this drug as best for the treatment of diabetes mellitus [45]. The prediabetic risk development in patients with impaired glucose tolerance in the Indian population has been significantly reduced with metformin [46]. American Diabetes Association and American Association of Clinical Endocrinologists recommend monotherapy with metformin for the earlier treatment of hyperglycaemia in adult T2DM patients [47]. Metformin is only biguanide currently in use, other derivatives like phenformin and buformin have been withdrawn from the market due to reported adverse effects like lactic acidosis and cardiac mortality in the early 1970s [48]. Metformin lowers blood glucose levels in both obese and non-obese T2DM patients and proves to be as effective as sulfonylurea derivatives [49]. Metformin
clinically reduces Hba1c level and can be used in combination with other oral antidiabetic drugs without gastrointestinal effects [50]. Other cellular mechanisms for metformin are via inactivation of gluconeogenesis enzymes production mediated through 5’AMP-activated protein kinase (AMPK), increase glucose uptake in skeletal muscles, and adipocytes [51]. AMPK mediated mechanism of action of metformin has been challenged by genetic loss-of-function studies and support by other AMPK-independent possible mechanisms that explain metabolic effects and side effects of metformin [52]. Metformin also acts through peroxisome proliferator-activated receptor (PPAR)-α, which induces insulin secretion and increases the glucagon-like peptide 1 (GLP-1) levels [53]. In skeletal muscles, metformin causes delocalization of GLUT1 protein but no effect on GLUT4 glucose transporters and stimulates glucose uptake [54]. Metformin has also been found to reduce cancer risk in T2DM [55]. Weight loss is associated with metformin in long term treatment. Hypoglycemia is usually not reported with metformin unless it is administered with sulfonylureas and insulin [56]. The gastrointestinal disturbances like anorexia, diarrhoea, metallic taste, and gut discomfort are common side effects reported with metformin but can be resolved with dose modifications [57].

**Sulfonylureas**

Sulfonylureas are developed in the 1920s and used in the treatment of T2DM. The chronic use of sulfonylurea normalizes insulin levels to baseline with reduced glucose levels. Sulfonylurea is a hormone-releasing agent [58]. Sulfonylurea acts as insulin secretagogues and stimulates insulin release from β-cells of the pancreas [59]. Sulfonylureas are classified into two group’s viz., first and second-generation agents. The less potent agents are including in first-generation agents, e.g., Acetohexamide, Chlorpropamide, Tolazamide, and Tolbutamide. The potent sulfonylureas are included in second-generation agents e.g., Glibenclamide, Glimepiride, Gliclazide, and Glipizide are available in the markets for the treatment of T2DM [60]. They inhibit ATP-sensitive K-channels in β-cell and initiate a cascade of events that result in the release of insulin from granules. ATP-sensitive K-channels is a complex consisting of two subunit proteins Kir6.2 (pore-forming subunit) and SUR1 (drug-binding regulatory subunit) [61]. SUR1 is a receptor binding site for sulfonylurea, which initiates the closing of ATP-K-channels and decreases efflux of potassium, which causes depolarization of the plasma membrane of β-cell. Cellular influx triggers depolarization and is mediated through voltage-dependent Ca2+-channels, and promotes the first and second phases of insulin secretion from preformed insulin granules [62]. Gliclazide and tolbutamide selectively inhibit ATP-sensitive K-channels in β-cell through SUR1 subunit, not SUR2 subunit in cardiac and smooth muscle. Other agents like glimepiride, Glimepiride, repaglinide, and meglitinide inhibit both subunits of ATP-sensitive K-channels [63]. Hypoglycaemia, an increase in body weights and secondary failure are noted side effects of sulfonylureas [64]. The chronic use of sulfonylureas significantly increases the risk of coronary heart disease in women [65].

**Meglitinides**

Meglitinides are nonsulfonylurea insulin secretagogues and act as rapid-acting prandial insulin releasers. They too increase insulin secretion similar to sulfonylurea. The two marketed meglitinides are repaglinide and nateglinide [64]. Meglitinides bind to Kir6.2 (pore-forming subunit) of ATP-sensitive K-channels present on the plasma membrane of β-cells of the pancreas [67]. Meglitinides are not effective in patients with dysfunction β-cells in the pancreas [68]. The monotherapy with repaglinide proves to be more effective in reducing Hba1c and fasting glucose levels in comparison to nateglinide [69]. Repaglinide shows better tolerance and appears to be a safe alternative in geriatrics diabetic patients [70]. They have a rapid and short duration of action due to short half-life and potentiates only first phase insulin release. This contributes to a lower risk of hypoglycaemia. The adverse reactions of meglitinides are similar to sulfonylurea but with a mild decrease in body weight [71].

**Thiazolidinediones**

Thiazolidinediones or glitazones are a new class of oral antidiabetic agents and are FDA approved drugs for T2DM, e.g., pioglitazone and rosiglitazone. Thiazolidinediones enhance insulin action, increase glucose muscle uptake, and cause suppression of gluconeogenesis. They reduce the concentration of circulating free fatty acids, triglycerides, and cause a rise in the levels of HDL and LDL cholesterol [72]. Thiazolidinediones decrease insulin resistance by repression of hepatic glucose output and increasing insulin-dependent glucose metabolism, thus, used in the treatment of T2DM associated with insulin resistance [73]. Thiazolidinediones enhance insulin sensitization and alter transcription of genes which modulate carbohydrate and lipid metabolism. The will ligands for peroxisome proliferative activated receptor-gamma (PPAR-gamma) in muscle, adipose tissue, and liver [71]. PPAR-gamma activation causes transcription in genes for lipoprotein enzyme lipase, adipocyte fatty acid-binding protein, fatty acyl-CoA synthase, glucokinase, and GLUT4. Thiazolidinediones diminish insulin resistance through the activation of endothelial signaling in skeletal muscle and liver [74]. Rosiglitazone and other PPAR gamma activator agents inhibit activation of the c-Jun NH-terminal kinase (JNK) signaling pathway and enhance the survival rate of β-cells of the pancreas [75]. The reported side effect of thiazolidinediones is oedema when used in combination therapy with insulin. Thiazolidinediones also aggravate the risk of congestive heart failure in T2DM patients [76].

**Alpha-glucosidase inhibitors (AGIs)**

Alpha-glucosidase is a membrane-bound enzyme present in the small intestine and assists in the absorption of carbohydrates after a meal [77]. Alpha-glucosidase acts through a mechanism of action, i.e., inhibition of carbohydrates absorption from the gastrointestinal tract. AGIs are pseudo-sugars that competitively and reversibly inhibit the α-glucosidase enzyme and cause a delay in the absorption of carbohydrates from the gut [78]. These are used in treatment of T2DM patients with uncontrolled hyperglycaemic, uncontrolled fasting glucose tolerance [79]. Voglibose, Miglitol, and Acarbose are the most widely used AGIs for the treatment of T2DM and decrease postprandial plasma glucose and insulin levels [80]. AGIs are most beneficial for T2DM patients having higher postprandial plasma glucose levels and normal HbA1C. A clinical situation where glucose levels are not controlled with monotherapy of other oral antidiabetic agents, diet, and exercise, AGIs can be used as first-line of a drug in combination therapy [78]. Acarbose is used as add-on therapy in poorly controlled T2DM patients with diet sulfonylurea and biguanide. The starting dose of AGI is low and administered after meals. These inhibitors reduce glucose level about 54 mg/dl and Hba1c 0.9% [81]. The common side effects of AGIs are flatulence, gut discomfort, diarrhoea, and bloating. Hepatic injury is a rare side effect reported with AGIs use [82].

**Dipeptidyl peptidase 4 inhibitor or gliptins**

There are two primary incretin hormones secreted from the intestine after post-meal are glucagon-like peptide-1 (GLP-1) and gastric inhibitory polypeptide (GIP). Incretin hormones stimulate insulin secretion from β-cells of the pancreas and regulate glucose homeostasis [83]. In response to elevated glucose concentration after a meal, incretin hormones (GLP-1 and GIP) enhance insulin release from β cells, and additionally GLP-1 decreases glucagon production [84]. In hypoglycaemia state, GIP promotes glucagon counter-regulation and increases α-cells sensitivity for glucose [85]. Incretin hormones action is mediated through activation of adenylyl cyclase and high level of cyclic adenosine monophosphate (cAMP) [86]. Incretin mimetics are a new class of drugs used in the treatment and management of T2DM, e.g., dipeptidyl peptidase 4 inhibitors (DPP-4 Inhibitor) and GLP-1 receptor agonist [87]. In plasma, dipeptidyl peptidase-4 (DPP-4) cause’s cleavage of two NH-terminal amino acids of both GIP and GLP-1, thus, block the action of incretin hormones which result in reducing insulin level and higher postprandial glucose [88]. DPP-4 Inhibitor effectively decreases Hba1c without causing an increase in the body weight [89]. Five DPP-4 inhibitors are available in market viz., sitagliptin, vildagliptin, saxagliptin, linagliptin, and alogliptin [90]. Sitagliptin is a selective and efficacious dipeptidyl peptidase 4 inhibitors used for the management of T2DM and well-tolerated with ongoing metformin therapy [91]. Recent studies reveal sitagliptin monotherapy reduces both fasting and non-fasting glucose level along with improved functioning of β cells in T2DM.
is nausea suppression, progressive stimulation of insulin secretion, increase in exendin and sulfonylurea for the management of T1DM. Clinical studies show that liraglutide in add-on therapy with metformin and thiazolidinediones decreases the higher glucose level and well-tolerated by T2DM patients [119]. Liraglutide significantly reduces fasting and non-fasting glucose levels with a lower incidence of hypoglycemia [120]. The use of these patients are at higher risk for the development of diabetes and cardiovascular disease. Such patients are advised for weight reduction. Liraglutide decreases insulin resistance, lowers blood pressure, and fatty acid concentration [121]. The obese patient with T1DM on insulin treatment, liraglutide is added for better control on glucose and glycated haemoglobin level. Liraglutide add-on therapy causes a reduction in insulin dose and body weight [122].

Albiglutide is a DPP-4 enzyme resistant long-acting GIP-1 receptor agonist and approved in many countries for the treatment of T2DM. It stimulates postprandial insulin release and delays gastric emptying resulting in reduced food intake [123]. Albiglutide administered once in a week through a subcutaneous route and exhibits less gastrointestinal side effects compared to other incretins mimic. It is the choice of agent in diabetes with renal failure [124]. Dulaglutide, a GIP-1 receptor agonist administered once in a week used as an adjunct to non-therapeutic methods and produced by recombinant DNA technology. Various clinical trial studies show dulaglutide monotherapy effectively control hyperglycaemia [125]. Dulaglutide efficiently reduce glycated haemoglobin with common side effect i.e. gastrointestinal discomfort [126]. Semaglutide is the latest approved GIP-1 receptor agonist for the treatment of T2DM by the US Food and Drug Administration and European Medicines Agency. It shows a significant decrease in glycated haemoglobin and systolic blood pressure along with the reduction of weight in T2DM patients [104]. Common side effects reported in saxagliptin monotherapy are diarrhoea, respiratory infection, mild hypoglycaemia, and headache [105]. Linagliptin, a xanthine base, is a selective inhibitor of the DPP 4 enzyme [106]. It decreases HbA1c levels in diabetes with a good tolerability profile in a patient with renal defects. It is well tolerated in obese, geriatric patients and does not require dose adjustment [107]. Linagliptin shows a better safety profile over sulfonylurea without causing an increase in body weight, stroke, and cardiovascular effects [108]. It enhances insulin sensitivity and reduces liver fat in hepatic steatosis [109]. Albiglutin is a highly selective noncovalent DPP-4 inhibitor and approved by the FDA in January 2013. It is prescribed for monotherapy or in combination with metformin and pioglitazone for T2DM [110]. Albiglutin is well tolerated and its monotherapy gives similar control over hyperglycaemia as other DPP-4 inhibitors without hypoglycaemia [111]. The combination therapy of albiglutin and pioglitazone improves β cell function while monotherapy of albiglutin gives better control of glycaemia [112]. A recent study shows that albiglutin prevents the progression of non-alcoholic fatty liver disease in T2DM patients [113]. The common side effects reported with albiglutin are respiratory infection, body pain, and nasopharyngitis [114].

**Incretin mimetic or GLP-1 receptor agonist**

Incretin mimetic or GLP-1 receptor agonists, a new generation of drugs that increase insulin release, inhibit glucagon secretion, and reduce appetite [115]. Seven incretin mimetics are available in market viz., Exenatide, Liraglutide, Albiglutin, Dulaglutide, Semaglutide, Taspoglutide and Lixisenatide. Exenatide (exenatide immediate-release injection, exenatide extended-release injection) is GLP-1 receptor agonist and the first member of incretin mimetic. It is approved by the FDA in 2005 as add-on therapy with metformin and sulfonylurea for the management of T2DM [115]. Exenatide and 39 amino acid peptide hormone having biological properties similar to glucagon-like peptide-1 (GLP-1). Exenatide is a synthetic product of exendin-4. Exenatide controls glycaemia via glucagon secretion suppression, progressive stimulation of insulin secretion, increase in β cell cellular mass, and enhancement of peripheral tissue glucose uptake [116]. The most common side effect reported with exenatide is nausea [117]. Liraglutide is a long-acting GIP-1 receptor agonist and having 97% resemblance to human incretin hormone, i.e., GLP-1 [118]. Liraglutide is an injectable GIP-1 analog approved by the FDA in February 2017 for the management of T2DM. The proposed mechanism of action of sitagliptin in type 1 diabetes mellitus mediated through suppression of glucagon production [90]. Vildagliptin is a new oral hypoglycaemic drug and acts as a selective inhibitor of the dipeptidyl peptidase-4 enzyme [93]. Vildagliptin causes prolonged inhibition of the DPP-4 enzyme and diminishes GLP-1 degradation. It increases insulin secretion, decreases glucagon concentration, inhibits gluconeogenesis, and enhances insulin sensitivity [94]. Vildagliptin increases insulin to glucagon ratio, which results in the enhancement of glucose sensitivity of α-cell and β-cell of the pancreas in T2DM patients. Insulin to glucagon ratio decreases in the hypoglycaemic state, resulting in the stimulation of hepatic glucose production [95]. A recent study shows that vildagliptin monotherapy or in combination with other antidiabetic agents is safe and well-tolerated but requires add-on therapy [96]. Vildagliptin and metformin in combination therapy efficiently lower blood glucose levels as compared to glimepiride-metformin treatment with a lower incidence of hypoglycaemia and a slight increase in body weight [97]. Saxagliptin is a reversible inhibitor of the DPP-4 enzyme, which binds covalently on catalytic active serine hydroxyl site and reduces degradation of incretin hormone and promotes insulin secretion [98]. DPP-4 enzyme inhibitors are used for better tolerability in the early stage of diabetes mellitus [99]. It is approved by the FDA in July 2009 and recommended an add-on to the non-therapeutic approach in T2DM, i.e., dietary modification and physical exercise [100]. Saxagliptin metabolite is less potent as a parent drug [101]. Saxagliptin is proved to be safe both in monotherapy and in combination. It is also effective in lowering glycylated haemoglobin, fasting plasma glucose, and non-fasting glucose [102]. Saxagliptin is found to be inferior to vildagliptin but superior to sitagliptin in controlling glucose levels [103]. Saxagliptin regularizes normal capillaries and ease homeostasis in T2DM [104]. Common side effects reported in saxagliptin monotherapy are diarrhoea, respiratory infection, mild hypoglycaemia, and headache [105]. Linagliptin, a xanthine base, is a selective inhibitor of the DPP 4 enzyme [106]. It decreases HbA1c levels in diabetes with a good tolerability profile in a patient with renal defects. It is well tolerated in obese, geriatric patients and does not require dose adjustment [107]. Linagliptin shows a better safety profile over sulfonylurea without causing an increase in body weight, stroke, and cardiovascular effects [108]. It enhances insulin sensitivity and reduces liver fat in hepatic steatosis [109]. Albiglutin is a highly selective noncovalent DPP-4 inhibitor and approved by the FDA in January 2013. It is prescribed for monotherapy or in combination with metformin or pioglitazone for T2DM [110]. Albiglutin is well tolerated and its monotherapy gives similar control over hyperglycaemia as other DPP-4 inhibitors without hypoglycaemia [111]. The combination therapy of albiglutin and pioglitazone improves β cell function while monotherapy of albiglutin gives better control of glycaemia [112]. A recent study shows that albiglutin prevents the progression of non-alcoholic fatty liver disease in T2DM patients [113]. The common side effects reported with albiglutin are respiratory infection, body pain, and nasopharyngitis [114].

**Sodium-glucose co-transporter-2 (SGLT2) inhibitors**

SGLT2 inhibitors offer a distinctive mechanism of action and consider the newer class of medicament in the treatment of Diabetes mellitus. SGLTs are active co-transporters that facilitate passive glucose transporters (GLUTs) for glucose reabsorption in the proximal convoluted tube. To date, six SGLTs were identified but SGLT1 and SGLT2 are considered valuable in glucose haemostasis [132]. SGLTs inhibition causes glucose excretion in urine, which results drop in blood glucose levels without the involvement of insulin. SGLT inhibitors such as dapagliflozin, canagliflozin, ertugliflozin, and empagliflozin having a low risk of hypoglycaemia and well-tolerated as monotherapy or in combinations other antidiabetic medications [133]. Dapagliflozin, a highly selective SGLT2 inhibitor, reduces blood glucose levels, glycosylated haemoglobin, and body weight with low-risk hypoglycaemia [134]. The common side effects of dapagliflozin in the clinical trial are female genital mycotic infections, urinary tract infections, and nasopharyngitis [135]. Canagliflozin is another SGLT2 inhibitor that effectively reduces plasma glucose levels with cardiovascular and renal benefits such as blood pressure and albuminuria [136]. It also reduces HbA1c in type 1 diabetic patients when used as an add-on to insulin therapy [137]. Ertugliflozin is a recent oral SGLT2 inhibitor that gets FDA approval as an antidiabetic drug. It prevents reabsorption of urine glucose and shows better control of body weight and blood pressure [138]. Ertugliflozin reduces cardiovascular risk due to a fair decline in blood pressure and body weight. Therefore, it is also recommended as an adjunct drug with non-therapeutic treatment methods like diet management and
exercise in T2DM [139]. Empagliflozin is a selective SGLT2 inhibitor, administered orally to decrease blood glucose levels via urinary glucose excretion. It is well tolerated and no dose adjustment is required in diabetic patients with renal or hepatic impairment [140]. Empagliflozin in a clinical trial on T2DM patients with cardiovascular disease showed marked reduction in cardiovascular risk but urosepsis or genital infection was sporadic reported [141].

Other agents

Amylinomimetic drugs are the synthetic analogue of amylin, a naturally occurring 37-amino-acid peptide hormone released from the pancreas, and helps in glucose homeostasis. T2DM patient possesses a lower level of amylin while T1DM patient does not produce amylin from the pancreas [142]. Pramlintide, an approved amylinomimetic drugs by the US FDA for the treatment of T1DM along with insulin, and in the case of T2DM, it is used as an adjunct with insulin, metformin and sulfonylurea [143]. Pramlintide reduces postprandial blood glucose in both T1DM and T2DM patients with side effects like hypoglycemia [144]. Bromocriptine mesylate, dopamine (D2) agonist approved by the FDA for the treatment of T2DM and used as both monotherapy and in combination with other oral hypoglycaemic agents. Bromocriptine resets the tone of dopaminergic and circadian neuronal activities in the hypothalamus and central nervous system [145]. The drug is well-tolerated, reduces glycated hemoglobin by 0.4-0.8%, and reduce cardiovascular risk [146].

Non-therapeutic methods

In recent decades, the incidence of diabetes is constantly increasing and widely spread among Indian communities especially in males [147, 148]. The higher prevalence of obesity and diabetes in the urban population is related to unhealthy lifestyles, i.e., lack of exercise, unhealthy dietary habits, and overeating [149, 150]. Diabetes mellitus even in early on stages cause a decline in skeletal muscle strength and loss of physical function [151]. Diet control and weight reduction have shown an advantageous effect on insulin action and weight gain. It is related in terms of change in macronutrients and abdominal fat [152]. In T1DM, variation in dose regimen of rapid and short-acting insulin has been made according to the carbohydrate content of meal for better control on hyperglycaemia [153]. A similar approach has been recommended for T2DM. In obese T2DM patients, increase in unsaturated fat consumption in meals causes a decrease in insulin sensitivity and an increase in body weight. Protein metabolism is altered in both types of diabetes mellitus. Type 1 diabetic patients tend to convert amino acids into glucose. Thus, modification in protein content has been recommended [154]. In T2DM, protein metabolism does not increase glucose levels instead protein ingestion causes stimulation of insulin release, C-peptide and glucagon [155]. Day to day lifestyle, lack of physical activities have increased developmental risk of T2DM in young, adult, and older people. Exercise or burning off extra calories found to be helpful in the prevention and control of T2DM and weight gain [156]. The eight-week-long exercise regime exhibit decrease in glycosylated haemoglobin irrespective to body weight. It is also reported that an increase in the intensity of physical exercise further results in the improvement in glycosylated haemoglobin [157]. Physical activities improve insulin sensitivity and decrease elevated glucose levels in the body. The risk of diabetes in a pre-diabetic patient with impaired glucose tolerance is reduced by a higher expenditure of energy and weight loss [158]. It also reduces the incidence of cardiovascular diseases associated with T2DM [159]. The patients counseling with informational booklet increase compliance for treatment and cause a decrease in glycaemic levels [160].

CONCLUSION

Diabetes mellitus, chronic condition with elevated glucose levels in the body. Its prevalence in the world increased rapidly, especially in urban and suburban areas. The insulin hormone maintains normal homeostasis in human beings. The management of insulin levels in terms of its concentrations and activities is the prime focus of current therapies. To achieve the afore-mentioned objective various medications i.e. insulin formulation and oral hypoglycaemic agents, were developed, which show a higher potency rate. A range of insulin preparation with the different onset of action was used to control glucose levels e.g., rapid-acting insulin, short-acting insulin, intermediate-acting insulin. Oral hypoglycaemic agents are used when insulin is present within the body but insulin sensitivity to a great extent reduced. These agents include metformin, sulfonylureas meglitinides, thiazolidinediones, DPP-4 inhibitors, GLP-1 receptor agonists, SGLT2 inhibitors. Metformin improves insulin response, sulfonylureas normalize insulin levels in the body, meglitinides stimulate insulin release, thiazolidinediones enhance insulin sensitivity, and DPP-4 inhibitors reduce glucose level without causing an alteration in body weight. GLP-1 receptor agonists lower the rate of glucose absorption from the intestine and SGLT2 inhibitors halts glucose reabsorption in the kidney. The changes in lifestyle i.e. exercise and diet restrictions, also help in maintaining glucose levels in the body.

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CONFLICT OF INTERESTS

There is no conflict of interest.

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