The Emergence of Sodium glucose Co-transporter 2 Inhibitors as a Successful Agent for Management of Diabetes Mellitus

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Abstract

Most of current guidelines for management of diabetes mellitus are recommending comprehensive strategy inclusive of adequate control of blood pressure and weight apart from hyperglycemia. Several innovative anti-diabetic molecules have been launched in recent years. Out of these agents, injectable incretin based therapy like GLP 1 receptor agonist has shown great promise, but high cost and injectable route of administration have prevented their wide acceptance. In the last 4 years, sodium glucose co-transporter inhibitor, which works in kidney to induce excess glucose elimination through urine, has been launched globally. This oral anti-diabetic agent works comprehensively and more intensively than existing dipeptyl peptidase 4 inhibitors (DPP4i). Apart from its excellent glycemic benefit with negligible risk of hypoglycemia, and extra-glycemic benefits like weight loss and blood pressure control, compatibility with almost all existing anti-diabetic agents, possible positive impact on cardiovascular and renal outcomes, and rapid reversal of glucotoxicity in type 2 diabetes mellitus patient whose renal function is intact, make them irresistible options for management of type 2 diabetes mellitus. However, despite huge advantages, injudicious use of agents this class throws up unique challenges like genito-urinary infections, eu-glycemic keto-acidosis, electrolyte imbalance etc.. This review tries to give a comprehensive and balanced view of this class of drug.

Keywords: Diabetes Mellitus Gliiflozin; SGLT2 inhibitors; Kidney; DKA

Introduction

Diabetes mellitus is a becoming a huge pharma-economic burden globally due to its substantial impact on increase in mortality, morbidity, and quality of life. The number of adults with diabetes in the world increased from 108 million in 1980 to 422 million in 2014. The diabetes prevalence rate has increased from 4.3% to 9.0% in male and 5.0% to 7.9% in females during the period from 1980 to 2014. The rate of increase in diabetes
prevalence is faster in low and middle income countries than higher income countries during last three decades. Out of the top five countries contributing half of total global population of patients with diabetes, three are from South-East Asian region (China, India and Indonesia), with a continuous increasing trend [1,2]. During the last three decades, several oral or injectable anti-diabetics molecules including some innovative ones like GLP 1 receptor agonist (GLP-1a), dipeptyl peptidase 4 inhibitors (DPP4i), newer generation of analogue insulin etc have evolved and incorporated into the armamentarium of diabetes management [3-5]. Diabesity, a term used to describe obesity amongst persons with diabetes has evolved as a major issue for clinicians as well as epidemiologist during last few decades [6]. Since 2012 onwards, SGLT2 inhibitors have been launched globally in different countries. From clinical perspective, the medical fraternity needs to evaluate whether this class of molecule deserves a respectable place in the treatment armamentarium to fight diabetes [7].

**SGLT2 Inhibitors – Mechanism of Action**

Kidneys play an important role in glucose homeostasis in body. Every day, they produce 15-55 gm of glucose by gluconeogenes is while simultaneously utilizing 25 gm. Along with this, kidneys also reabsorb 180 gm of glucose per day at proximal convoluted tubules (PCT) [8]. This renal glucose reabsorption is happening mainly through SGLT 1 and SGLT2 co-transporters. Almost 90% of glucose re-absorption is by SGLT 2 co-transporter situated at S1 segment of PCT and rest 10% glucose re-absorption is by SGLT1 co-transporter situated at S3 segment of PCT (Figure1) [9].

![Figure 1: Structure of nephron and precise location and primary functions of Sodium Glucose co Transporters (SGLT). Adopted and modified from: Vlotides and Mertens Nephrol Dial Transplant.](image)

In type 2 diabetes mellitus (T2DM), over expression of SGLT2 and SGLT1 co-transporters lead to enhanced renal threshold for glucose upto ~250 mg/dl which in normal subject without diabetes is ~ 180 mg/dl (10, 11) SGLT2 inhibitors block reabsorption of glucose by SGLT2 co-transporter resulting in excretion of ~70 gm/day of in production in liver by stimulating pancreatic alpha cells to secrete glucagon, improve insulin sensitivity and glucose utilization in skeletal muscles. Thus SGLT2 inhibitors counterbalance multiple aspects of diabetes urin which is equivalent to approximately ~280 kcal/day. This is accompanied by a loss of ~400 ml of water per day due to osmotic diuresis [11,12]. In addition to this, they improve pancreatic beta-cell function indirectly by reducing glucotoxicity, increase lipid oxidation in adipose tissues, increase endogenous glucose pathophysiology, except perhaps increase in endogenous glucose production [11].
Glycemic and Extra-Glycemic Benefit with SGLT2 Inhibitors

In the last 6 years, almost six SGLT inhibitors (canagliflozin, dapagliflozin, empagliflozin, tofogliflozin, loseogliflozin, ipragliflozin) have been launched globally [13-17]. There is no head to head trial looking at comparative efficacy amongst different SGLT2 inhibitors. Most widely used SGLT2 inhibitors are canagliflozin, dapagliflozin and empagliflozin (not in any order). Few meta-analyses involving different agents within the class of SGLT2 inhibitors show canagliflozin 300 mg as the most efficacious vs all others in terms of glycemic efficacy and extra-glycaemic benefit like weight loss but with higher incidences of adverse events such as genito-urinary infection (Table 1). On the other hand dapagliflozin 10 mg and empagliflozin 25 mg both show balance in terms of safety and efficacy as per meta-analysis within SGLT2 inhibitors class [18].

<table>
<thead>
<tr>
<th>Name of SGLT2 inhibitor</th>
<th>HbA1c Changes (%)</th>
<th>Weight Reduction (in Kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Canagliflozin 300 mg</td>
<td>-0.9 CI (-1.0,-0.8)</td>
<td>-2.5 CI (-2.8,-2.1)</td>
</tr>
<tr>
<td>Canagliflozin 100 mg</td>
<td>-0.8 CI (-0.9,-0.7)</td>
<td>-1.9 CI (-2.2,-1.5)</td>
</tr>
<tr>
<td>Empagliflozin 25 mg</td>
<td>-0.7 CI (-0.8,-0.6)</td>
<td>-2.3 CI (-2.6,-1.9)</td>
</tr>
<tr>
<td>Dapagliflozin 10 mg</td>
<td>-0.7 CI (-0.7,-0.6)</td>
<td>-2.1 CI (-2.5,-1.8)</td>
</tr>
<tr>
<td>Empagliflozin 10 mg</td>
<td>-0.6 CI (-0.7,-0.5)</td>
<td>-2.2 CI (-2.5,-1.9)</td>
</tr>
<tr>
<td>Dapagliflozin 5 mg</td>
<td>-0.6 CI (-0.7,-0.4)</td>
<td>-1.6 CI (-2.0,-1.2)</td>
</tr>
</tbody>
</table>

Table 1: Salient efficacy features of different SGLT2 inhibitors from multitudes of clinical trials. 18 SGLT2: Sodium Glucose Co-Transporter2; HbA1c: Glycosylated haemoglobin.

The clinical implications of benefits of SGLT2 inhibitors transcend beyond hyperglycaemia control because of their remarkable ability in reducing blood pressure and weight. There is approximately ~2kg to ~2.5kg placebo corrected weight reduction when SGLT2 inhibitors are added on to metformin therapy for 2 years, an effect that is sustained over 4 years [19-21]. More importantly this class of drug was found to be associated with 2/3rd fat mass loss and only 1/3rd of lean mass loss which resulted in reduction of waist circumference [22]. SGLT2 inhibitors also address the core modifiable challenges of Asian Indian phenotypic characteristics by improving glucose disposal rate in skeletal muscles, improving beta cell function, reducing serum triglycerides, reducing leptin etc. [23-25]. Three other SGLT2 inhibitors namely tofogliflozin, loseogliflozin and ipragliflozin developed in Japan have shown similar glycaemic and extra-glycaemic benefits [13-15].

SGLT2 Inhibitors and Incretin Therapy

The first decade of this millennium witnessed a paradigm shift in the approach to the management of T2DM with the advent of incretin based therapy, namely dipeptyl peptidase 4 inhibitors (DPP4i) and glucagon like polypeptide 1 receptor agonist (GLP-1RA). DPP4i has seen wide acceptance globally as add on to metformin therapy, or even as accompaniment to insulin. But the same cannot be said about GLP-1Ra due probably to high cost especially in developing countries, inspite its’ potent glycemic as well as extra-glycemic benefit like weight loss which is an important component of comprehensive anti-diabetic management as suggested by many of current guidelines (Table 2) [26-30].

<table>
<thead>
<tr>
<th>Guidelines</th>
<th>Glycaemic control</th>
<th>Weight (kg)</th>
<th>Blood pressure</th>
</tr>
</thead>
<tbody>
<tr>
<td>WHO 200626</td>
<td>&lt;7.0%</td>
<td>&lt;25 (M) &lt;24 (F)</td>
<td>&lt;94 (&lt;37) Male &lt;130/80</td>
</tr>
<tr>
<td>EASD-ESC 200727</td>
<td>≤6.5%</td>
<td>&lt;25 (M)</td>
<td>&lt;80 (&lt;32) Female &lt;130/80</td>
</tr>
<tr>
<td>ADA 201228</td>
<td>&lt;7.0%</td>
<td>≤25 (M)</td>
<td>&lt;80 (&lt;35) Female &lt;130/80</td>
</tr>
<tr>
<td>ADA 201129</td>
<td>≤6.5%</td>
<td>-</td>
<td>&lt;80 (&lt;35) Female &lt;130/80</td>
</tr>
<tr>
<td>CDA 200830</td>
<td>≤7.0%</td>
<td>&lt;25 (M)</td>
<td>&lt;80 (&lt;35) Female &lt;130/80</td>
</tr>
</tbody>
</table>

Comparison of results of meta-analysis of SGLT 2 inhibitors and DPP4i indicates superiority of SGLT2 inhibitors over DPP4i in terms of efficacy and extra-glycemic benefits with similar overall safety. Like incretin based therapy, SGLT2 inhibitors also protect against gluco-toxicity-induced apoptosis of pancreatic β-cells which is remarkable [31,32].

**SGLT2 inhibitors: advantages**

SGLT2 inhibitors work through insulin independent mode of action [33]. In addition to this, their efficacy also depends on serum glucose level, due to which chances of developing hypoglycaemia with their usage is negligible [34]. SGLT2 co-transporters are expressed also at alpha cells in pancreas and inhibition of this co-transporter leads to slight increase of serum glucagon level within physiological range that resulted into mild increase endogenous glucose production in liver [11,35]. SGLT2 inhibitor can be used as1st line agent in mono-therapy in patients who cannot tolerate metformin. By and large this new class of drug has seen widespread use as an add on as second or even third line of anti-diabetic agent to metformin, sulphonyl urea, DPP4i, thiazolidinedione etc.SGLT2 inhibitor can be used as1st line agent in mono-therapy in patients who cannot tolerate metformin. It can be safely used concomitantly with insulin or GLP1a. Such flexibility and compatibility would go a long way in achieving comprehensive diabetes management outcome [33].

**Certain challenges SGLT2 inhibitors – myths and reality**

Every good thing may come with certain challenges and SGLT2 inhibitors are no exceptions. However it is important to delve deeply and take a holistic view in this regard.

**Urinary Tract infection or Genital mycotic infection.**

Patients with diabetes are at risk of developing bacterial or fungal infection. These infections are frequently associated with uncontrolled hyperglycaemia. Lower urinary tract infection (UTI), which is common in type 2 diabetes patients has been reported in up to 22% of such patients [36]. Infection risk is directly related to glycaemic status with the risk becoming approximately 2-fold greater (adjusted RR, 1.76; 95% CI, 1.30–2.38) for patients with HbA1c levels in the highest quintile despite on-going anti-hyperglycaemic therapy [37]. Apart from bacterial infections, mycotic infection in diabetes is quite common with uncontrolled hyperglycaemia. Risk of developing vulvo-vaginal infection is 81% greater in females with T2DM (adjusted relative risk [RR], 1.81; 95% CI, 1.64–2.00) [37]. Mycotic infections are definitely higher with the use of SGLT2 inhibitors, mainly because of glucosuria. However, most of such events are mild to moderate in intensity responding promptly to initial course of anti-fungal therapy and once uncontrolled hyperglycaemia gets corrected, the rate of recurrence is less [38,39].

**Hyponatremia or Electrolyte Imbalance**

Due to the unique mechanism of action, SGLT2 inhibitors blocks sodium glucose co-transporter 2 in proximal convoluted tubule in nephron preventing glucose and sodium (Na+) entry into the tubular epithelial cells. Apart from this SGLT pathway, there are several other pathways responsible for sodium reabsorption in nephron. For example, sodium -hydrogen anti-port system plays major role in sodium reabsorption. Although 60-70% of sodium reabsorption is happening through proximal convoluted tubule, just <5% of that would be mediated by the sodium glucose co-transporter SGLT2 (Figure 1) [40]. No wonder that in 13 placebo controlled trial involving Dapagliflozin, (phase 2b and 3), mean change of serum and potassium from baseline over 24 weeks in the Dapagliflozin arm was similar with the placebo arm [40, 41].

**Diabetic Keto-Acidosis (DKA)**

Risk of DKA in patients with type 2 diabetes on SGLT2 inhibitors is quite low, probably numbering less than one in 1000 to one in 10,000. [42] There are two types of diabetic keto-acidosis, one with significant hyperglycaemia, and another with euglycaemia or moderate hyperglycaemia (blood glucose<300 mg/dl). The later type is popularly known as eu-glycaemic diabetic keto-acidosis (eu-DKA). This can happen in a background of too less amount of insulin secretion leading to restriction in the usage of glucose as substrate to generate energy. T his triggers exaggerated breakdown of fat to produce free fatty acid to be utilised for ketogenesis and finally to enter into process of energy production through Kreb’s cycle. Certain steps are critical to avoid DKA while prescribing SGLT2 inhibitors. If the patient is on insulin, the insulin dose can be reduced as per demand but should not be withdrawn totally. It is mainly for this reason, this new drug should not be
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prescribed in type 1 diabetes patient till enough clinical data showing its safety are available (Table 3) [43].

<table>
<thead>
<tr>
<th>Dos</th>
<th>Don'ts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reduce Fluid loss</td>
<td>Don't use in acute illness</td>
</tr>
<tr>
<td>Encourage adequate rehydration</td>
<td>Don't stop ongoing insulin totally</td>
</tr>
<tr>
<td>Encourage adequate carbohydrate intake</td>
<td>Don't use in Type 1 diabetes mellitus</td>
</tr>
<tr>
<td>Stop SGLT2 inhibitors &gt;48hours before surgery</td>
<td></td>
</tr>
<tr>
<td>Be on alert for early signs and symptoms of DKA</td>
<td></td>
</tr>
</tbody>
</table>

Table 3: Measures to prevent diabetes ketoacidosis (DKA) in SGLT2 inhibitor treated patients. Some of the precautionary measures are widely practiced while prescribing metformin.

**Acute Kidney Injury (AKI)**

In May 2016, FDA has come up with acute renal failure warning against canagliflozin and dapagliflozin based on 101 cases reported at FDA adverse effect reporting system during the period from March 2013 to October 2015. These cases were out of 1.5 million SGLT2 inhibitor users. Out of these, 73 cases were due to Canagliflozin and 28 cases due to dapagliflozin. No case was reported for empagliflozin. Such differential reporting is obviously due to the fact that in the US, canagliflozin was launched in March 2013 followed by dapagliflozin in January 2014 and empagliflozin in September 2014 [44-47]. Of the 101 cases, 51 reported concomitant ACE inhibitor use, 26 reported concomitant diuretic uses, and 6 reported concomitant non-steroidal anti-inflammatory drug (NSAID) use. A prior history of chronic kidney disease was reported in 10 of the 101 cases. In some cases, dehydration or hypotension was reported [44].

Regarding renal safety with SGLT2 inhibitors, in phase 2b/3 trial, eGFR data is available with canagliflozin, dapagliflozin, and empagliflozin. For canagliflozin, 2 years data, for empagliflozin 1 year data and with canagliflozin both 2 and 4 years data of eGFR are available. In Empa-REG trial, 4 years eGFR data is available with empagliflozin. Cutting across all the trials with SGLT2 inhibitors, though there were initial drop of eGFR within 1-2 weeks of initiation of treatment, normalization in the subsequent weeks on continuous usage of the product was a general rule rather than exception [21, 48-51]. The drug safety communication stated that health care professionals should carefully consider certain factors, including state of dehydration; chronic kidney insufficiency; congestive heart failure; and concomitant medications such as diuretics, angiotensin-converting enzyme inhibitors (ACEI), angiotensin receptor blockers (ARBs), and nonsteroidal anti-inflammatory drugs (NSAIDs) [44]. Beyond these precautionary steps, these drugs are considered by and large safe from renal point of view.

**Fracture**

After anecdotal reports and some clinical trial data evaluation, in September 2015, USFDA enhanced the level of warning for canagliflozin indicating increased risk of bone fractures, and added new information about decreased bone mineral density [52]. The occurrence of bone fractures was evaluated in nine pooled clinical trials with a mean duration of exposure to canagliflozin of 85 weeks. The incidence rates of adjudicated bone fractures were 1.1, 1.4, and 1.5 per 100 patient-years of exposure in the comparator (placebo or active), canagliflozin 100 mg, and canagliflozin 300 mg groups, respectively. Fractures were observed as early as 12 weeks after treatment initiation, and were more likely to be precipitated by low intensity trauma (e.g., arising after falls from no more than standing height) and affect the upper extremities [52]. There is no significant alteration of bone turnover markers or bone mineral density with dapagliflozin, in men as well as postmenopausal women with T2DM after 50 weeks and 102 weeks of follow-up [22,53]. No increase in fractures was observed in pooled data from patients with normal to mildly impaired renal function across the dapagliflozin clinical studies [44]. Fracture incidence was comparable in both empagliflozin and placebo arm (3.9% vs 3.8%) in phase 2b/3 f trials [47]. Overall, incidence of fracture is about 3 per 1000 patients treated with SGLT2 inhibitors [46,51,54].

**Hypovolemia or Dehydration**

SGLT2 inhibitor works as osmotic diuretic and due to its mechanism of action there could be fluid loss of ~400 ml/day which is roughly equivalent to just one extra void of urine. This urine output is spread throughout the day, and does not cause nocturia [11]. However, in climatic condition of tropical countries like the Indian subcontinent, especially in the background of excessive heat exhaustion, this may present realistic health challenges such as drug induced hypovolemia and electrolyte imbalance. Counseling must include advice to maintain adequate fluid and electrolyte intake, and awareness about symptoms of dyselectrolytemia. One
should also ensure that SGLT2 inhibitors are not co-prescribed with loop diuretics. However, there is no need to monitor serum electrolytes or renal function tests in persons on long term SGLT2 inhibitors treatment [55].

Safety and efficacy are important considerations during selection of any molecule, and if these are important than even a pioneer molecule like metformin is also having safety issue like lactic acidosis, sulphonylurea is having issues like hypoglycaemia, weight gain, so also other popular anti-diabetic and anti-hypertensives. It is of paramount importance on the part of the clinicians to understand the balance between safety and efficacy, and that is where right selection of patient for SGLT2 inhibitor plays an important role [56,57].

**Best Sglt2 Inhibitor Within Class**

Differentiation within the class of SGLT2 inhibitors is inconclusive. First of all, so far, there is no head to head trial comparing the efficacy. Regarding safety analysis, especially for cardiovascular events, only Empa-Reg trial of empagliflozin has been published so far. This trial has shown that empagliflozin can significantly reduce cardiovascular death, possibly through haemodynamic benefit. However Empa-Regenrolled patients with T2DM of whom more than 99% had prior cardiovascular event. The ongoing CANVAS trial for canagliflozin and DECLARE-Timi 58 trial for dapagliflozin have enrolled varying number of T2DM patient not only with prior history of prior cardiovascular event (secondary prevention), but also subjects at risk of cardiovascular disease (primary prevention). The later cohort is a true representative of vast majority of patients encountered in day to day practice. Meanwhile, meta-analysis of cardiovascular outcome of phase 2b/3 trials of both canagliflozin and dapagliflozin has shown trends similar to that of Empa-Reg trial. Overall, such data dispel the doubt regarding cardiovascular safety of SGLT2 inhibitors and make them strong contender for a place in anti diabetic armamentarium at various stages of the disease [45,46,58,59].

**Right Selection Of Patient For Sglt2 Inhibitor**

SGLT2 inhibitor works through insulin independent pathway and this can be prescribed across the entire disease spectrum of T2DM, as suggested by most of the guidelines. It can be used as add on to metformin, sulphonylurea, insulin etc as second or third line agent. Few situations where it should not be used are patients with estimated glomerular filtration rate (eGFR) below 45 ml/min (), type 1 diabetes, concomitant acute illness or surgery, severe dehydration, pregnant ladies, reproductive age female planning pregnancy, very frail elderly patient or patient with history of recurrent infection. It can also be considered across the wide spectrum of body mass index (BMI) as glycaemic benefit result is similar irrespective of baseline BMI [60-62].

**Conclusion**

SGLT2 inhibitors belong to a class of oral anti-diabetic agent which actually works as poly-pill. Its pleotropic umbrella covers excellent control of hyperglycaemia, along with satisfactory control of hypertension, and reasonable reduction of body weight, each of which is an important objective of comprehensive anti-diabetic management on its own merit. In large clinical trials, there are strong early signals of cardiovascular and renal risk reduction which, it is hoped, will be supported by high quality evidences in the coming years. SGLT2 inhibitors should be considered as an early add to standard anti-diabetic therapy in appropriate T2DM patient.

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**Conflict of interest**

All the authors have contributed independently to express their personal views in this paper and there is no conflict of interest.

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