Management of CV risk in T2DM: The beginning of a new era

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\textbf{Introduction}

Individuals with type 2 diabetes mellitus (T2DM) experience increased morbidity and mortality secondary to vascular damage involving both the microvasculature [retinopathy, nephropathy, neuropathy] and macrovasculature (myocardial infarction, stroke, amputation) \cite{1}. Diabetes is the leading cause of blindness and end stage renal disease in western countries \cite{2}, and T2DM individuals have a 2.5-4 fold increase in the risk of cardiovascular disease \cite{3}. Cross sectional studies have demonstrated that one-third to one-half of all people with diabetes have evidence for organ damage \cite{4}. Although not everyone with diabetes is destined to develop complications, a recent epidemiological study \cite{4} reported that two or more complications are apparent in almost one-fifth of people with diabetes.

\textbf{Diabetes and Microvascular Complications}

Two landmark studies, the Diabetes Control and Complications Trial (DCCT) \cite{5} and United Kingdom Prospective Diabetes Study (UKPDS) \cite{6}, as well as others \cite{7-10}, unequivocally have documented that improved glycemic control in T2DM subjects reduces the risk of microvascular complications, demonstrating that chronic exposure of tissues to hyperglycemia triggers pathologic processes that lead to eye, kidney and nerve damage. A 1% decrease in HbA1C has been shown to reduce the incidence of microvascular complications by \(-35\%\) \cite{5,6,9} and maintaining the HbA1C \(<6\%\) has been reported to completely prevent the development of retinopathy in T2DM subjects \cite{8}. Further, in the DCCT continuation study (EDIC), subjects who initially were treated aggressively with intensive insulin therapy and achieved a HbA1C \(<7\%\) maintained a lower risk for microvascular complications many years later, even though their HbA1C increased to 8\% compared to subjects who initially were less well controlled \cite{11}. Similar results have been reported in the long term extension of UKPDS \cite{12}. Insulin was utilized to achieve intensive glycemic control in the DCCT and insulin, sulfonylurea and merformin were utilized in the UKPDS to lower the plasma glucose concentration. Conversely, improved glycemic control in T2DM patients with long-standing poor glycemic control has less of an impact to reduce the risk of microvascular complications \cite{13}. More recent studies have demonstrated that newer antidiabetic agents, e.g. GLP-1 receptor agonists (GLP-1 RAs), SGLT2 inhibitors, and pioglitazone, which are very effective in lowering the plasma glucose concentration, also reduce the risk of diabetic microvascular complications. Collectively, these observations emphasize the importance of achieving good glycemic control at the time of diagnosis \cite{14} and maintaining the level of glycemic control, independent of therapeutic strategy utilized, to reduce the risk of microvascular complications. The results of these studies have prompted the professional health care organizations like American Diabetes
Association [15], American College of Physicians [16], American Association for Clinical Endocrinology (AACE) [17], the European Association for the Study of Diabetes [15] and the Canadian Diabetes Association [18] to recommend glycemic goals as close to normal as possible while avoiding hypoglycemia in newly diagnosed T2DM patients without clinically manifest vascular disease. The ADA’s target goal for HbA1C is ≤ 7.0%, while the AACE’s and the EASD’s goal is ≤ 6.5%.

Macrovascular Complications in T2DM

Subjects with T2DM have a markedly increased risk of cardiovascular disease [myocardial infarction and stroke] and a worse prognosis following any cardiovascular event [3,19]. Further, T2DM increases the risk of heart failure in the absence of ischemic heart disease, and the presence of T2DM worsens the prognosis in patients with heart failure [20]. Although hyperglycemia is an important risk factor for microvascular complications, improved glycemic control only modestly reduces or has no effect to reduce the increased risk of cardiovascular events [6,13,21,22]. UKPDS demonstrated that hyperglycemia per se has a less prominent role in the development of macrovascular complications compared to microvascular complications [6]. Further, in UKPDS [23] and VADT [24] it took more than 10 years to observe a modest CV benefit following improved glycemic control. Most T2DM individuals manifest moderate - severe insulin resistance which is associated with multiple metabolic abnormalities (obesity, dyslipidemia, hypertension, endothelial dysfunction, procoagulant state), all of which are important risk factors of CVD [25]. This cluster of cardiovascular/metabolic disturbances is known as the Insulin Resistance (Metabolic) Syndrome and likely contributes to the increased CVD risk in T2DM. Hypertension [26,27] and dyslipidemia [28] are major risk factors for coronary artery disease and many studies have documented that reduction of blood pressure and correction of dyslipidemia significantly reduces CVD. Nonetheless, despite reduction of blood pressure [27,28] and plasma LDL cholesterol to target levels [28], cardiovascular risk in T2DM subjects remains greater than in nondiabetic subjects [29,30]. Many recent studies [31-39] have demonstrated that insulin resistance, independent of the associated metabolic abnormalities, is an important risk factor for cardiovascular disease. Although hyperglycemia is only a weak risk factor for cardiovascular disease, although recent clinical outcome trials [37-43] have demonstrated that members of SGLT2 inhibitor class, GLP-1 receptor agonists (GLP-1 RA) and pioglitazone significantly reduce CVD risk in T2DM patients in whom traditional CV risk factors (e.g. blood pressure, dyslipidemia) are optimally controlled. In the following discussion, we will briefly review the results of these studies and discuss the implications of these findings for the care of patients with T2DM.

Insulin Sensitizers and CVD

Because insulin resistance is closely linked with CVD risk in T2DM, it follows that improving insulin sensitivity with an insulin sensitizers would reduce CV risk, independent of their glucose lowering action. Pioglitazone is the only true insulin sensitizer available for treatment of T2DM [44-46]. In addition to lowering plasma glucose concentration, pioglitazone decreases insulin resistance (by 35-40%) in skeletal muscle and liver [47] decreases plasma triglyceride concentration, increases HDL cholesterol, converts small dense atherogenic LDL particles to larger more buoyant ones, and reduces blood pressure [48,49]. Pioglitazone also reduces plasma FFA, adipocytokines/other inflammatory markers/procoagulant factors, and increases plasma adiponectin [44-48], all of which would be expected to provide cardiovascular benefit. Thus, pioglitazone would be expected to provide additional cardiovascular benefits, independent of the reduction in plasma glucose concentration [50,51]. Consistent with this, pioglitazone has been shown to slow the progression in carotid intima medial thickness in the Chicago [52] and ACT NOW studies [53], and to reduce coronary atheroma volume [54] in subjects with type 2 diabetes in the Periscope study. In large clinical outcome studies, pioglitazone significantly lowered the incidence of 3-point MACE [non-fatal myocardial infarction, non-fatal stroke, CV death] in T2DM patients. In PROactive [41], 5238 T2DM patients with existing CVD were treated for 2.9 years with pioglitazone or placebo plus standard of care for glycemic control and CV risk factors. 3-point MACE,
the main secondary endpoint, was significantly reduced by 16% (HR=0.84, p=0.027). In IRIS [40], 3876 insulin resistant (HOMA-IR >3.0), nondiabetic, insulin resistant individuals with recent [within 6 months] ischemic stroke or TIA were randomized to pioglitazone or placebo for 4.8 years. Pioglitazone caused a 24% reduction in fatal/nonfatal stroke plus myocardial infarction (HR=0.76, P=0.007). Because glycemic control in subjects in the placebo arm in the PROActive study were treated to target and participants in IRIS study did not have T2DM, the results of these studies demonstrate that pioglitazone reduced CVD risk independent of its glucose lowering action.

GLP-1 RAs and CVD risk in T2DM

Multiple mechanisms contribute to the glucose lowering action of GLP-1 RA, e.g. enhanced beta cell function, inhibition of plasma glucagon concentration, delayed gastric emptying, and suppression of hepatic glucose production. GLP-1 RAs also suppress appetite, resulting in significant weight loss and, indirectly, improving insulin sensitivity [55]. GLP-1 RAs also decrease plasma triglyceride and increases plasma HDL concentrations and reduce blood pressure [56]. These later actions of GLP-1 RA would be expected to have a favorable effect to reduce cardiovascular risk [56]. Three recent large outcome trials, LEADER [37], SUSTAIN [38] and EXSCEL [43] have examined the effect of once daily liraglutide, once weekly semaglutide and once weekly exenatide, respectively on cardiovascular risk (3-point MACE: non-fatal myocardial infarction, non-fatal stroke and CV death) in T2DM patients. All 3 studies primarily recruited T2DM patients with established CVD. In LEADER [37], 9340 T2DM patients (82% with prior CV event) were randomized to liraglutide, 1.8 mg/day, or placebo for a mean of 3.8 years. Investigators were blinded to the study intervention and instructed to maintain HbA1c <7.0% with any antidiabetic medication except GLP-1 RA or DPP4 inhibitor. Compared to placebo, liraglutide caused a 13% reduction in 3-point MACE. In SUSTAIN [38] 3297 T2DM patients (83% with established CVD) were randomized to semaglutide, 0.5 and 1 mg/day, or placebo and followed for 2 years. Similar to LEADER, investigators were instructed to lower HbA1c to <7.0% according to local guidelines without using incretin-based therapies. Semaglutide caused a 26% reduction in the primary outcome (3-point MACE) which fell just short of statistical significant (p=0.06) but discontinuation [-25%] of exenatide was very high in the EXSCEL study. Two aspects of LEADER, SUSTAIN and ESXCEL deserve emphasis: (i) patients at higher CVD risk benefited more from GLP-1 RA treatment. In a combined analysis of the three long acting GLP-1 RAs, the risk of CVD in subjects with T2DM and established CVD (secondary CV prevention) was significantly reduced (by 13%) in subjects receiving GLP-1 RA (liraglutide, semaglutide and exenatide) (HR=0.87, 95% CI=0.81-0.93, p=0.001). Conversely, subjects with T2DM without established CVD (primary CVD prevention) did not benefit from GLP-1 RA therapy (HR=1.07, 95% CI=0.88-1.3, p=0.49). (ii) the benefit of liraglutide, semaglutide and exenatide was evident on top of optimal control of traditional CV risk factors [37,38,43].

SGLT2 Inhibitors and CVD risk

Sodium-glucose cotransporter 2 (SGLT2) inhibitors have a unique mechanism of action [57]. They lower the plasma glucose concentration by inhibiting renal glucose reuptake and produce glucosuria. This unique mechanism of action, in addition to lowering plasma glucose concentration, results in multiple other metabolic and hemodynamic actions which could favorably benefit CVD risk in T2DM patients [57-61]. SGLT2 inhibitors cause (i) weight loss of 2-3 kg; (ii) reduction of ~5 mmHg in systolic blood pressure; (iii) osmotic diuresis which results in a modest decrease in extracellular volume of ~ 5-10%; (iv) increase in total body fat oxidation and ketone production;
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(v) a small increase in plasma HDL cholesterol and decrease in plasma triglycerides; (vi) improvement in total body insulin sensitivity (17%). Because these metabolic actions of SGLT2i could improve CVD risk, one might expect that this class of drugs would lower CVD risk in T2DM, independent of their glucose lowering effect. Two large clinical outcome studies have examined the effect of empagliflozin and canagliflozin on 3-point MACE in T2DM patients. In EMPA REG OUTCOME trial [39], empagliflozin caused a 14% reduction (P=0.04) in 3-point MACE in 7020 T2DM patients with established cardiovascular disease over 3.1 years and this was driven by a robust 38% reduction in CV mortality. Further, the reduction in 3-point MACE was accompanied by a 39% reduction in the rate of hospitalization for heart failure.

One surprising finding in EMPA-REG, unlike LEADER, SUSTAIN, EXSCEL, IRIS and PROactive, separation between the empagliflozin and placebo curves occurred very early, such that reduction in the primary outcome was evident at 3 months after starting treatment. In CANVAS/CANVAS-R [42], 10,142 T2DM patients (66% with established CVD) were randomized to receive canagliflozin or placebo for 3.6 years. The reduction in 3-point MACE caused by canagliflozin was identical to that observed with empagliflozin (14%, p<0.001). Similar to EMPA-REG, the separation between the two curves occurred early after starting therapy and was accompanied by a 33% reduction in the rate of hospitalization for heart failure. Collectively, the results of CANVAS and EMPA-REG OUTCOME studies suggest that: (i) it is likely that the CV benefits of canagliflozin and empagliflozin are class effect; (ii) although the mechanisms underlying the CV benefits of both drugs are not completely understood [62], it is likely their mechanism of action to reduce MACE is not due to inhibition of atherosclerosis. (iii) the rapid onset of CV benefit, and the marked reduction in hospitalization for heart failure suggest that the hemodynamic actions of SGLT2 inhibitors play an important role in their CV protective effect; (iv) similar to the LEADER, SUSTAIN, EXSCEL, IRIS and PROactive studies, only subjects with established CVD benefited from the treatment, while T2DM without existing CVD did not benefit from SGLT2 inhibitor therapy; (v) similar to GLP-1 RAs and pioglitazone, the CV benefit of SGLT2 inhibitors was observed in subjects with optimal treatment of traditional CVD risk factors, e.g. blood pressure, LDL and aspirin treatment.

**Implication or Care**

Overwhelming evidence supports that early glycemic control, regardless of the therapeutic strategy utilized, reduces the risk of microvascular complication in T2DM patients. However, the results of recent clinical outcome trials demonstrate that antidiabetic agents vary in their effect on CVD risk in T2DM (FIGURE 1) and we are entering a new era in T2DM management [63]. Despite improved glycemic control, DPP-4 inhibitors and basal insulin failed to reduce CV risk in T2DM patients [64-67], while GLP-1 RAs, SGLT2 inhibitors, and pioglitazone significantly reduce both plasma glucose concentration and CV risk (3-point MACE). Further, the CV benefit of GLP-1 RAs, SGLT2 inhibitors and pioglitazone is independent of their glucose lowering effect and occurs on top of optimal treatment of traditional CV risk factors. Further, the beneficial CV action of GLP-1 RAs, SGLT2
inhibitors and pioglitazone was evident only in T2DM patients with established CVD. Thus, GLP-1 RA, SGLT2 inhibitors and pioglitazone, not other antihyperglycemic agents, will reduce both microvascular and macrovascular complications in this group of T2DM patients. Therefore, evidence-based medicine dictates that in newly diagnosed T2DM patients with established CVD and in long-standing T2DM patients who experience a CV event, GLP-1 RAs, SGLT2 inhibitors and pioglitazone should be favored over other antidiabetic agents. On the other hand, in T2DM patients without established CVD, emphasis should be placed on lowering the HbA1c to target regardless of the antihyperglycemic strategy utilized.

References


