LETTER TO THE EDITOR

SGLT-2 Inhibitors: Fighting Cardiovascular Disease Mortality in Diabetes

Sir,

Cardiovascular (CV) events are the leading cause of mortality in diabetic patients. Managing this macro-vascular complication has been the forefront concern of both endocrinologists and cardiologists. Since the UK Prospective Diabetes study showed that better glycemic control improves CV outcomes in newly diagnosed diabetes patients,1 what followed from here was an immense number of trials trying to find the best possible outcome for such patients. However, in early 2008, ACCORD (Action to Control Cardiovascular Risk in Diabetes) study debunked this aggressive approach to maintain a tighter glycemic control, as it may lead to an increased risk of death due to hypoglycemia.2

The hastened approval of TZDs by the FDA to treat diabetes mellitus (DM) patients further accelerated this calamity in terms of increased deaths due to heart failure and myocardial infarction (MI). It provoked them to design trials in such a way as to lower the glycemic index without compromising the CV outcome. In the past decade, we have seen novel drugs entering the realm of good glycemic control, increased rate of HbA1c reduction, obesity-treatment and CV and renoprotective qualities. These new drug classes are the glucagon like peptide-1 (GLP-1) agonists and the sodium-glucose-linked-transporter-2 (SGLT-2) inhibitors.

The CANVAS (Canagliflozin Cardiovascular Assessment Study) programme compared efficacy of canagliflozin with placebo in diabetics who had an elevated risk of CV disease. The primary outcome of the study was focused on a combination of CV death, nonfatal MI, or nonfatal stroke.

The trial concluded a significant decrease in the incidence of the primary outcome event in the canagliflozin group: 26.9 vs. 31.5 participants with an event per 1000 patient-years (hazard ratio, 0.86; 95% CI, 0.75 to 0.97; p<0.001 for noninferiority; p=0.02 for superiority). The only serious adverse event seen in the group of patients taking canagliflozin was a higher risk of amputation of toes, feet, or legs; and was observed more in patients who had a history of peripheral vascular disease or a previous amputation.3

The CANVAS-R trial was a subset of the CANVAS programme and had the primary objective to study the effect of canagliflozin on kidney disease progression, with similar study design to the CANVAS trial.4 It concluded that progression of albuminuria occurred less frequently with canagliflozin.3 This finding can possibly be attributed to the nephro-protective effects of SGLT-2 inhibitors. By increasing angiotensin, they can restore the intraglomerular pressure, thus demonstrating vasodilatory and anti-inflammatory properties.5

These findings from the CANVAS programme have played a significant role in evolving our understanding of the effects of canagliflozin, and the broader SGLT2 inhibitor class, on a variety of efficacy and safety outcomes of crucial standing to patients with diabetes.4

The SGLT-2 class is coming up to be the front runner in managing micro- and macro-vascular complications along with good glycemic control. Due to its non-insulin effect, it can be paired up with insulin or other insulin secretagogues to further improve HbA1c. These are new horizons in fighting secondary complications of diabetes. Further trials are anticipated which should focus on all drugs of this class with a homogenous inclusion criteria of patients, similar risk factors, and a standardised outcome, addressing differences in race, pre-existing vascular, cardiac and renal conditions, and a longer duration of follow-up to obtain the most accurate results.

REFERENCES


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