Glycemic Control and Cardiovascular Outcomes in Diabetes
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Abstract
While type 1 Diabetes Mellitus (DM) is characterized by insulin deficiency due to pancreatic beta cell destruction, type 2 DM is characterized by a state of long standing insulin resistance (IR), compensatory hyperinsulinemia and varying degrees of elevated plasma glucose (PG), associated with clustering of cardiovascular (CV) risk and development of macrovascular disease prior to diagnosis of DM. Coronary artery disease (CAD) accounts for 70% of mortality and morbidity in patients with diabetes.

Studies made in diabetes care have helped prevent or reduce microvascular complications in type 1 and 2 diabetes. However the same cannot be said about macrovascular disease.

Despite all data concerning the association of diabetes and cardiovascular disease (CVD), the exact mechanism by which diabetes is linked to atherosclerosis is incompletely understood, this is especially true in case of hyperglycemia. The positive effect of intensive glucose management in comparison to non intensive glucose control is far from proven.

DCCT and UKPDS studies have shown that while a glycemic control is important for reaching long term macrovascular complications, early glucose control is far more rewarding (metabolic memory). Later trials like ACCORD, ADVANCE and VADT don’t advocate tight glycemic control. In fact, ACCORD trial has shown increased mortality with tight glucose control.

Tight glucose control may be beneficial in selected patients with short disease duration, long life expectancy and no CVD. In critically ill patients a blood glucose target of 140-180 mg% is fairly reasonable and achievable.

The ESC/EASD guidelines of October 2013, like those of ADA, AHA and ACC continue to endorse a treatment target for glucose control in diabetes of HbA1c <7%, based predominantly on microvascular disease with acknowledged uncertainty regarding the effect of the intensive glucose control on CVD risk.

Management of hyperglycemia in diabetics should not be considered in isolation; diabetics require multifactorial intervention for hypertension, dyslipidaemia and microalbuminuria besides hyperglycemia. In fact combined use of antihypertensives, aspirin and lipid lowering agent makes it difficult to discern salutary effects of anti hyperglycemic therapy.

Keywords: Diabetes; Glycemic control; Hyperglycemia

Introduction
Diabetes mellitus (DM) is a condition defined by an elevated level of blood glucose. Type 1 diabetes is characterized by deficiency of insulin due to progressive destruction of pancreatic beta cells, progressing to absolute insulin deficiency. Type 2 diabetes is a combination of insulin resistance and beta cell failure in association with obesity and sedentary life style. However, not all overweight/ obese individuals have diabetes and vice versa.

The increase prevalence of diabetes worldwide has led to a situation where approximately 360 millions people had diabetes in 2011, of which 95% would have type 2 DM [1]. This number is estimated to increase to 552 million by 2013 and it is presumed that half of these will be unaware of their diabetes status.

The prevalence of diabetes is increasing world wide and more people with diabetes will die or be disabled as a consequence of vascular complications. Prospective studies have shown unambiguous association of blood glucose and glycated hemoglobin level with the risk of major cardiovascular events. In case of subjects with type 1 diabetes, in spite of the fact that CV rate is significantly lower compared with population with type 2 diabetes, their relative risk for coronary heart mortality is 7 fold higher than in matched counterpart without disease.

In spite of all these data concerning the association of diabetes and cardiovascular diseases (CVD), the exact mechanism by which diabetes is linked to atherosclerosis remains incompletely understood. This is especially true in case of hyperglycemia. The role of non-glycemic factors accompanying vast majority of patients with diabetes such as hypertension, dyslipidaemia and hemorrheological abnormality are better understood and appear to be independent of hyperglycemia. There also has been data regarding the future impacts of statins, aspirin, ACE inhibitors and aggressive control of blood pressure on progression of CV disease. In contrast, the positive effect of intensive glucose management on CV disease outcome is far from proven. Even some studies show a negative influence. The objective of the present article is to analyze trials related to glycemic control in diabetics and assess its impact on CV outcomes.

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Glycemic Continuum and CVD

Type 2 DM develops following a prolonged period of euglycemic insulin resistance (IR) which progresses with development of beta cell failure to frank diabetes with increase risk of vascular complications. While microvascular complications like retinopathy, nephropathy and neuropathy develop with onset hyperglycemia, macrovascular complications like coronary artery disease, cerebrovascular disease and peripheral arterial disease (PAD) appear earlier during the stage of IFG and IGT. Thus these complications are already established when type 2 DM is diagnosed. Over 60% pts with type 2 DM develop CVD which is a more severe and costly complication than retinopathy.

Molecular Basis of CVD

Insulin resistance has an important role in the pathophysiology of diabetes and CVD. Both genetic and environmental factors facilitate its development. The development of CVD in people with IR is characterized by early endothelial dysfunction and vascular inflammation leading to monocyte recruitment, foam cell formation and subsequent development of fatty streaks [1]. Over many years this leads to atherosclerotic plaque which in presence of enhanced inflammation becomes unstable and rupture to promote occlusive thrombus production. Atheroma from people of diabetes has more lipid, inflammatory change and thrombus than those free from DM. These changes occur over a 20-30 year period and are mirrored by the molecular abnormalities seen in untreated insulin resistance and DM.

Type 2 DM patients are obese and the release of free fatty acids (FFA) & cytokines from adipose tissue directly impairs insulin sensitivity in skeletal muscle and adipose tissue. FFA induces reactive oxygen species production, blunts activation of IRS 1 and PI3K – AKT signaling leading to down regulation of insulin responsive GLUT 4 (Figure 1).

Hyperglycemia decreases nitric oxide bioavailability and affects vascular function involving over production of reactive oxygen species [1]. The mitochondrial electron transport chain is one of the first targets of high glucose with a direct increase in super oxide anion formation. A further increase in super oxide anion formation is driven by a vicious cycle involving ROS induced activation of PKC [1]. Mitochondrial ROS in turn activates cascades involved in the pathogenesis of the CV complications including polyol flux, AGE and RAGE. Hyperglycaemia induced ROS generation is involved in the persistence of vascular dysfunction despite normalization of blood glucose levels. This phenomenon is called metabolic memory which explains why vascular complication progresses despite intensive glycemic control. Elevated ROS generation despite euglycemic sensitivity undermines the clinical gold standard of indexing type 2 efficacy by blood glucose status.

Insulin resistant macrophage increases expression of oxidized LDL scavenger receptor-B, promoting foam cell formation and atherosclerosis. Macrophage dysfunction provide a crucial link between diabetes and CVD by both enhancing it and by contributing to the development of fatty streaks and vascular damage.

Impact of Glucose Control on CVD and its Complications

Randomised controlled trials provide compelling evidence that microvascular complications of DM are reduced by tight glycemic control. However the same cannot be said about macrovascular disease. Several prospective trials have been conducted which have so far failed to provide any conclusive evidence of the superiority of glycemic control in reducing macrovascular complications, or death rates in people with advance disease or those with long duration of diabetes.

Long term effect of glycemic control

A. Diabetes Control and Complications Trial (DCCT) and Epidemiology of Dibetes Interventions and Complications (EDIC)

In DCCT the rate of CV events was not significantly altered in the intensive treatment group of patients with type 1 DM. After termination of study, 93% of the cohort were followed for additional 11 years under EDIC, during which the differences in HbA1C disappeared. During the combined 17 years follow up, the risk of any CV event was reduced significantly in the intensive group by 42% (9.63% p<0.01) [2].

B. United Kingdom Prospective Diabetes Study: (UKPDS)

C. In the UKPDS trial, 3867 newly diagnosed subjects with type 2 DM were randomised to an intensive glucose control arm involving use of sulfonylurea or insulin and a conventional arm employing life style management. A subgroup of over weight subjects were included in the study that compared intensive glucose control with metformin (n=343) against conventional therapy (n=411). In the insulin and sulfonylurea group, a mean HbA1C level of 7% was achieved versus 7.9% in the control arm over 10 years. Intensive control decreased risk for a composite end point of all diabetes related complications (RRR=12%,p=0.029),and significantly improved microvascular disease risk (RRR=25%, p=0.01), where as a trend towards decreased risk of MI was observed with intensive control (14.8% vs. 16.8%, p=0.052, statistically not significant). Stroke was
numerically increased (5.6% vs. 5.2%, p=0.05). In over weight subjects, metformin had better glucose control (Alc >7.4% vs. 8%) as well as significantly improved risk for MI (RRR=39%, p=0.01) and for all cause mortality (RRR=26% p=0.011).

In extension phase UKPDS study, the patients were followed up for additional 10 years after completion of the trial, during which difference between HbA1c levels in both the groups disappeared. The follow up showed significantly reduced risk for MI in those originally randomised to intensive glycemic control both in insulin and sulfonylureas groups (RRR=15%, p<0.01) and in the metformin group (RRR=33%, p=0.05) [3].

There was also significantly 13% reduction in all cause mortality in the intensively treated group. This persistent benefit generated from early strict glycemic control is known as legacy effect, which outlives the original reduction of HbA1c and subsequent loss of glycemic control. These observations are similar to those seen in DCCT follow up EDIC study where CV events, non fatal MI, stroke and CV death were reduced by 57% despite loss of glycemic separation [2].

**Combined UKPDs and DCCT / EDIC study show that**

1. Glycemic control is important for reducing long term macrovascular complications.
2. Very long follow up period is necessary to demonstrate any benefit.
3. Early glucose control is important (metabolic memory).

**Medium term effects of glycemic control**

**Action to Control Cardiovascular Risk in Diabetes (ACCORD):**
The landmark study was designed to determine whether CV disease event rate could be reduced by intensively treating hyperglycemic hypertensive and dyslipidemia in a double 2x2 factorial design. The trial was based on the hypothesis that a 1.5% difference in HbA1c would result in 15% difference in a population of high risk diabetic individuals having a 3% annual CVD event rate [4].

The study included 10,251 patients with established type 2 DM and 1/3rd having a CV event. Patients were randomised to intensive glucose (targeting HbA1c < 6% and achieving a level of 6.4%) or standard therapy (targeting HbA1c of 7.0 - 7.9 % and achieving level of 7.5%). A variety of glucose lowering therapy was used. There was non significant trend towards reduction in primary outcome of trial (a composite of non fatal MI, stroke or CV death) with intensive control. However, unexpectedly there was higher all cause mortality (CR-1.22, 95 % CI-1.01-1.46, p=0.04). Higher rate of severe hypoglycaemia and weight gain were reported in intensive glycemic control group. Patients with high HbA1c level at base line were at higher rate of hypoglycaemia as were those who did not respond properly with a fall of HbA1c in intensive control group. The explanation for incremental mortality remains unresolved; possible explanations include hypoglycaemia precipitating CV death, pernicious effects of specific drug or combinations and a chance finding.

**Advance trial:** The study was conducted to determine whether intensive lowering would reduce risk of microvascular and macrovascular events in individuals with type 2 DM and vascular risk factors compared to standard conventional case. The study involved 11,140 subjects. The mean duration of follow up was 5 years. The patients were randomised to intensive versus standard glucose control with gliclazide plus other drugs in the intensive arm compared with other drugs in the standard control group. Mean HbA1c achieved was 6.5% in the intensive group compared with 7% in the standard group. The incidence of combined major macrovascular and microvascular events was significantly reduced (HR=0.9, 95% CI 0.82-0.98, p=0.01) in the intensive control group. This was primarily driven by reduction in progression of albuminuria or emergence of new nephropathy. The CV component of the primary event was not significantly reduced by intensive glucose control. There was no evidence of increase in all cause mortality. Actually there is a non significant trend towards reduction in all cause mortality [5].

**The VADT study:** The trial included American veterans, and 90% were males. A variety of glucose lowering agents was used including metformin, glimepiride, rosiglitazone and insulin. An HbA1c of 6.9% was achieved in the intensified control arm compared with HbA1c of 8.4% in the standard treatment arm. After a median follow up of 6.5 years, no significant lowering of composite CV outcomes was noted in the intensive control group. Severe hypoglycaemia was more prevalent in the intensive control group. Benefits of intensive control were apparent only in individuals with shorter duration of diabetes, lower HbA1c and absence of CVD at base line. Table 1 shows the baseline characteristics of ACCORD, ADVANCE & VADT trials [6].

**Insights from ACCORD, ADVANCE and VADT Trial**

- Important finding of all 3 studies is the suggestion that a beneficial effect of glycemic control intervention is more likely in association with less disease duration.
- In the ACCORD study participants with base line A1c < 8% rather than having adverse effects of intensive glycemic treatment on mortality, showed a significant reduction in primary outcome favoring such treatment. Similarly in ADVANCE trial, the combined macro and microvascular primary outcome benefit of glycemic control intervention was seen in participants without a baseline history of macrovascular disease. Similarly in the VADT trial, patients who had composite outcome event had longer diabetes duration, higher HbA1c and coronary arterial calcification.
- Effect of hypoglycaemia may be of importance. In the ACCORD study, although investigators stated that this was not a mediator of increased mortality associated with intensive therapy, intensive interventions was associated with significant severe hypoglycaemia. The ADVANCE and VADT study group similarly have reported high incidence of severe hypoglycaemia.
- A meta analysis of these 3 trials suggest that HbA1c reduction of 1% is associated with 15% of relative risk reduction in non fatal MI, but without benefit on stroke or all cause mortality.
- Conclusion from these 3 trials is that intensive glycemic control should be appropriately applied in an individualized manner taking into account age, duration of diabetes and history of CVD.
- Despite the fact that ACCORD, ADVANCE and VADT showed no benefit of intensive glucose control on primary CV endpoints in Type 2 DM, subgroup analyses suggest that any potential benefit on CV outcomes and mortality depends upon multiple interrelated factors such that medications capable of exerting direct CV therapeutic effects may be required to see a CV benefit.
One should also remember that HbA1c cutoff makes less sense for the cardiac events because cardiovascular risk depends upon various strong risk factors like hypertension and smoking [7] (Table 1).

**Glucose Control in ACS**

Elevated plasma glucose during an ACS is associated with a serious prognosis in patients with DM than without diabetes. Hyperglycemia may relate to previously undetected glucose perturbations but also to stress induced catecholamine release increasing FFA concentration, decreased insulin production and increasing insulin resistance and glycolgenolysis with a negative impact on myocardial metabolism and function.

Two strategies have been tried in an attempt to improve prognosis in patients with ACS

**Metabolic modulation**

Metabolic modulation by means of glucose-insulin-potassium infusion regardless of presence of DM or elevated PG, is based on the assumption that increase in intracellular potassium stabilizes the cardiac myocytes and facilitates glucose transport into the cell. Other potential benefits include decreased production of FFA, improved use of glucose for energy production and improved endothelial function and fibrinolysis. Despite these proposed mechanistic benefits of glucose, potassium and insulin therapy, the strategy has been proven futile in CREATE trial which enrolled more than 20000 patients with ACS. The NICE SUGAR trial in fact demonstrated a lack of effect may be due to increased PG or negative effect of fluid load induced by G&K therapy. This lack of effect may be due to increased PG or negative effect of fluid load induced by G&K therapy.

The DIGAMI trial, which is often misinterpreted as a trial of intensive glucose control is actually a glucose insulin infusion therapy trial [8]. The first DIGAMI trial randomised 620 patients with DM and AMI to >24 hrs insulin-glucose infusions followed by multi-dose insulin, or routine glucose lowering therapy. Mortality after 3-4 yrs was significantly reduced in the intervention group [8]. However DIGAMI-2 failed to replicate this prognostic benefit. The plausible reason for this discrepancy was that in DIGAMI-1 admission HbAlc decreased more (1.5%) from a higher level (9.1%) compared with 0.5% from 8.3% in DIGAMI-2. Since DIGAMI-2 failed to replicate this prognostic benefit. The plausible reason for this discrepancy was that in DIGAMI-1 admission HbAlc decreased more (1.5%) from a higher level (9.1%) compared with 0.5% from 8.3% in DIGAMI-2. Since DIGAMI-2 trial did not achieve a difference in glucose control between intensively treated and control groups, it is still an open question as to whether glucose lowering is beneficial.

**Glucose control in ICU setting**

In 2001 Van den Berghe published a randomised controlled trial of critically ill surgical pts showing that tight glucose control reduced hospital mortality [9]. Since the greatest decrease in death occurred in subgroup of pts with multi system organ failure, it was speculated that benefits of tight glucose control might extend to medical ICU patients as well. However subsequent trials by the same group couldn’t demonstrate any benefit with tight glycemic control. Further recent trial like VISEP and European glucontrol showed trend for increased rate with tight glucose control. The NICE SUGAR trial in fact demonstrated an actual 14% increase in mortality rate with intensive glucose regimen [10].

Few of these trials assessing glucose control in ICU settings included ACS patients. Therefore, general applicability of the observation remains uncertain. Because of paucity of data on tight glycemic control a glucose target of <180 mg/dl is a reasonable approach in ACS pts.

**Why Lower is not Necessarily Better?**

The UKPDS study was the first to provide evidence that in newly diagnosed type 2 DM patients intensive glucose control may reduce the risk of microvascular complications, also with modest effect on CV outcomes. Thus the concept ‘the lower, the better’ (glucose level) was proposed by all diabetology guidelines as a paradigm for type 2 DM patients. However, this concept has been challenged by 3 landmark trials: ACCORD, ADVANCE and VADT.

Numerous potential reasons have been put forth to explain the lack of benefits with intensive glucose control therapy. These include pernicious effects of specific drugs or drug combinations, increased incidence of hypoglycaemia precipitating CV death and a mere chance finding. The current glycemic target is <7% of HbA1c with individualization of therapy (Table 2).

**Hypoglycemia and Adverse CV Events**

In the ACCORD trial, which included diabetic patients with CV

|                | ACCORD | ADVANCE | VADT |
|----------------|--------|---------|------|
| N              | 10,251 | 11,140  | 1791 |
| Age (mean, years) | 62    | 66      | 60   |
| BMI (mean, kg/m²) | 32    | 28      | 31   |
| Follow-up (mean, years) | 3.5  | 5       | 5.6  |
| A1c target     | <6.0% versus 7.0%-7.9% | ≤6.5% versus “standard” | <6% versus 8%-9% |
| Baseline A1c (mean) | 8.3% | 7.5%    | 9.4% |
| Endpoint A1c (mean) | Intensive 6.4% Standard 7.5% | Intensive 6.43% Standard 7.0% | Intensive 6.9% Standard 6.9% |
| Severe hypoglycemic events | Intensive 10.5% Standard 3.5% | Intensive 2.7% Standard 1.5% | Intensive 8.5% Standard 8.5% |
| Weight change   | Intensive +3.5 kg Standard +0.4 kg | Intensive −0.1 kg Standard −1.0 kg | Intensive +8.1 kg Standard +4.1 kg |
| Major macrovascular or microvascular event | Not reported | 0.9 (0.82-0.98), P = 0.01 | 0.88 (0.74-1.05), P = 0.14 |
| Nonfatal MI/stroke, CV death | HR 0.9 (0.78-1.04), P = 0.16 | 0.94 (0.84-1.06), P = 0.32 | Not reported |
| All-cause mortality | HR 1.22 (1.01-1.46), P = 0.04 | 0.93 (0.83-1.06), P = 0.28 | 1.07 (0.81-1.42), P = 0.62 |
| Nonfatal MI | HR 0.76 (0.62-0.92), P = 0.004 | 0.98 (0.77-1.22), P = NS | 0.82 (0.59-1.14), P = 0.24 |

ACCORD: Action to Control Cardiovascular Risk in Diabetes trial; ADVANCE: Action in Diabetes and Vascular Disease: preterAx and diamicron N-MR Controlled Evaluation trial; A1c: Glycosylated Hemoglobin; BMI: Body Mass Index; CV: Cardiovascular; MI: Myocardial Infarction; VADT: Veterans Affairs Diabetes Trial

Table 1: Baseline characteristics of ACCORD, ADVANCE & VADT trials.
While an HbA1c target of less than 7% to reduce microvascular disease is a generally accepted level, the evidence for an HbA1c target in relation to macrovascular risk is less compelling.

Consensus indicates that an HbA1c of less than 7% should be targeted but with acknowledgement of need to pay attention to the individual requirement of the patient.

Fasting plasma glucose should be less than 120mg % (7.2mmol/l) and postprandial less than 160-180mg % (9-10mmol/l) on an individualized basis. Ideally tight glycaemic control should be started early in the course of the disease in younger people and without attendant comorbidities.

Stringent targets like HbA1c 6-6.5% may be considered in selected patients with short disease duration, long life expectancy & no CVD, if it can be achieved without hypoglycaemia or other adverse effect.

For critically ill indoor patients insulin therapy is indicated at a threshold of no greater than 180mg % (10mmol/l) (ADA2008).

Once insulin therapy has been started in critically ill patients a glucose range of 140-180 mg% is recommended.

With the preferred method of intravenous insulin infusion, frequent glucose monitoring is essential to minimize occurrence of hypoglycaemia and to achieve optimal glucose control.

 Tight glucose control (80 – 110 mg %) has not been associated with mortality benefit in many trials. In past some trials show increase mortality.

**Table 2: Current Glycemic Targets (ESC/EASD guidelines)** [1]

| Glycemic Target | Value |
|-----------------|-------|
| 1. Fasting glucose | 70-110 mg/dL |
| 2. Postprandial glucose | <140 mg/dL |
| 3. HbA1c | <7% |
| 4. Hemoglobin | <120 mg/dL |
| 5. Creatinine | <1.5 mg/dL |
| 6. Urea | <20 mg/dL |
| 7. Serum creatinine | <1.5 mg/dL |
| 8. Serum potassium | 3.5-5.0 meq/L |
| 9. Serum sodium | 135-145 meq/L |

**Cardiovascular Effects of Drugs used in Diabetes**

Few data are available regarding the net cardiovascular safety and efficacy of medications used to control glucose level in diabetes.

Metformin has best track record of safety, tolerability and low hypoglycaemia risk. This drug remains the drug of first choice.

Concern always exists regarding ability of sulfonylurea, to impair ischaemic preconditioning. However, UKPDS has been able to allay such fear to some extent.

Of thiazolidinediones, rosiglitazone was withdrawn from market because of fear of increased myocardial infarction risk. Recently, it has been reintroduced. Pioglitazone reduces myocardial infarction risk but can cause fluid retention.

Dipeptidyl peptidase 4 inhibitors have so far shown to have no adverse cardiovascular outcomes. Their safety track appears good.

Insulin increases the risk of hypoglycaemia and retrospective studies show adverse outcome when insulin is used in diabetics with heart failure.

**Conclusions**

As the disease diabetes assumes alarming proportions and threatens to become the modern pandemic, every effort should be made to prevent diabetes related cardiovascular complications. Interventions to reduce fasting blood glucose levels have unfortunately been not translated to better cardiovascular outcomes in all individuals. Recent trials like ACCORD, ADVANCE, and VADT challenge this proposition. However, metaanalysis of these trials suggests that the subgroups of diabetics with shorter duration of illness are beneficial from tight glycemic control (HbA1c<7%). Hypoglycaemia is always an issue, when the physician aims for tight glycaemic control. As hypoglycaemia adversely affects cardiovascular homeostasis, every effort should be made to avoid it at all costs while going for tight glycaemic control.

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