# How low should you go? New data on intensive 

 glucose lowering in type 2 diabetes
## Tony O'Sullivan


#### Abstract

In the pages of this journal, Colin Kenny discussed the disappointing outcome of three long-term trials of intensive glucose-lowering in type 2 diabetes with regard to cardiovascular disease events (Kenny, 2008). These trials (Veterans Affairs Diabetes Trial, ADVANCE [Action in Diabetes and Vascular disease: preterAx and diamicroN mr Controlled Evaluation] and ACCORD [Action to Control CardiOvascular Risk in Diabetes]) were presented and discussed at the American Diabetes Association's 68 ${ }^{\text {th }}$ Scientific Sessions in June 2008. This article summarises the new information and seeks answers to the practical dilemmas arising from their confounding results.


Ever since the UKPDS (UK Prospective Diabetes Study) hinted at a possible reduction in cardiovascular risk, at least among overweight people on metformin (UKPDS Study Group, 1983), the search for how to limit the excess cardiovascular disease (CVD) morbidity and mortality associated with type 2 diabetes has been on. The multifactorial approach has become standard care for type 2 diabetes with treatment targets tightening considerably over time; and recently attention has returned to glycaemic control as a factor. Perhaps driven by the growing understanding of diabetes' pathophysiology and the increasing range of oral agents and insulin preparations which make tighter glycaemic control achievable, studies have sought to clarify whether normalizing $\mathrm{HbA}_{1 \mathrm{c}}$ levels below those achieved in the UKPDS might have a greater impact on the remaining excess CVD risk. The results of these trials are now emerging, and are not only disappointing but also raise new concerns about intensive blood glucose control. This article summarizes what we now know, and what it might mean day-to-day for people
with diabetes and clinicians.

The Veterans Affairs Diabetes Trial The Veterans Affairs Diabetes Trial was a longterm ( 7.5 years) trial among older veterans, most of whom were male (see American Diabetes Association [2008] for more information). The aim was to assess the impact on cardiovascular disease (CVD) events related to intensive blood glucose control in addition to optimal control of associated CVD risk factors; the control group were provided standard blood glucose control. Participants had a mean $\mathrm{HbA}_{1 c}$ of $9.5 \%$ at randomisation, half had abnormal lipids, $80 \%$ had hypertension and over $40 \%$ had prior CV events.

The first aim of achieving and maintaining CVD risk factor control was achieved in both groups. Blood pressure (BP) and low-density lipoprotein cholesterol levels fell to a mean of $127 / 70 \mathrm{mmHg}$ and $2.0 \mathrm{mmol} / \mathrm{L}$, respectively. Percentage of participants smoking also fell from $16 \%$ to $10 \%$ across both groups. Target $\mathrm{HbA}_{1 \mathrm{c}}$ was between $8 \%$ and $9 \%$ in the control group (a median of $8.4 \%$ achieved), and as near

Article points

1. Three recent long-term trials have investigated intensive glucoselowering in patients with type 2 diabetes.
2. The results reveal a higher mortality rate among those patients subject to intense glucose control than in those with more relaxed glucose targets and less medical attention.
3. In this article, we try to come to grips with the clinical implications of these unexpected results.

Key words
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- Cardiovascular disease

Tony O'Sullivan is a GP at the Irishtown Health Centre, Dublin.
normal as possible in the intervention group (a median of $6.9 \%$ achieved). Insulin was used in $90 \%$ of the intensive group after the first year, and $74 \%$ of the control group. Severe hypoglycaemia was much higher in the intensive group ( $21 \%$ of participants versus $10 \%$ of controls).

Over a mean 6.25 years' follow up, CV events were $30 \%$ lower than expected in both groups. There were 231 events in the intensive group, and 263 in the control group, representing around a quarter of each group. The difference did not reach significance. The microvascular outcomes will be presented at the European Association for the Study of Diabetes Annual Meeting in September 2008.

## The ADVANCE trial

The ADVANCE (Action in Diabetes and Vascular disease: preterAx and diamicroN mr Controlled Evaluation) trial was a factorial randomised controlled trial with BP and blood glucose lowering arms. The BP arm was reported on in 2007 (ADVANCE Collaborative Group, 2007). The glycaemia arm randomised approximately 11000 people with type 2 diabetes to intensive or routine blood glucose lowering (ADVANCE Collaborative Group, 2008). Participants were on average 66 years of age, with a mean diabetes duration of 8 years. While not selected for added CV risk factors, around one third had experienced a previous CV event. Mean $\mathrm{HbA}_{1 c}$ at randomisation was $7.5 \%$.

The intensive blood glucose lowering arm involved the use of gliclazide modified release $30-120 \mathrm{mg}$ daily, otherwise glucose-lowering therapy was at the clinician's discretion. Over the trial period, each patient in the intensively treated group visited their health professional an average of 31 times (compared with 11 times each for those in the standard group). By the end of the study $40 \%$ were using insulin compared with $24 \%$ of the control group.
The ADVANCE trial was powered to detect a $16 \%$ difference in primary outcomes over 3.5 years, but the actual rates were low; so, in order to increase the power of the study, the glucose arm was extended to 5 years. Key results are as shown in Table 1. The primary end points were composites of major macroand microvascular events, assessed both together and separately.
Achieving a reduction in $\mathrm{HbA}_{1 c}$ of 0.7 percentage points over a median of 5 years reduced nephropathy by $21 \%$ and microvascular events by $14 \%$. The absolute risk reduction for microvascular events was $1.5 \%$, giving a number needed to treat (NNT) of 67. A little marginal perhaps, but at least this clarifies the benefit that can be achieved among typical patients with type 2 diabetes at this $\mathrm{HbA}_{1 \mathrm{c}}$ range.
In this author's opinion this trial firmly establishes the lack of benefit from intensifying glucose control when it comes to CV reduction. While those participants with pre-existing CVD had

| Outcome | Intensive control | Standard control | RR | $P$-value |
| :---: | :---: | :---: | :---: | :---: |
| Final $\mathrm{HbA}_{1 \mathrm{c}}$ (\%) | 6.5 | 7.3 | - | - |
| Combined micro and macrovascular events (\%) | 18.1* | $20^{*}$ | 0.9 | 0.01 |
| Microvascular events (\%) | 9.4 | 10.9 | 0.86 | 0.006 |
| New or worsening nephropathy (\%) | 4.1 | 5.2 | 0.79 | 0.006 |
| New or worsening retinopathy (\%) | 6.0 | 6.3 | 0.95 | 0.5** |
| Macrovascular events (\%) | 10.0 | 10.6 | 0.94 | 0.32** |
| Cardiovascular death (\%) | 4.5 | 5.2 | 0.88 | $0.12{ }^{* *}$ |
| All-cause mortality (\%) | 8.9 | 9.6 | 0.93 | 0.28** |

## Page points

1. The ACCORD study set out to address the advantages or otherwise of tight glycaemic and blood pressure control in a range lower than those currently recommended, with HbA1c targeted below 6\%.
2. The glucose-lowering arm of the ACCORD study was halted in February 2008, due to $22 \%$ more patients in the intensiveglucose control arm having died than those with more relaxed blood.
3. Twice as many patients in the intensive group as in the standard group gained $>10 \mathrm{~kg}$ of weight during the ACCORD study period, which might have led to a divergence in risk.
4. ACCORD participants who experienced a serious hypoglycaemica were much more likely to die, regardless of which treatment group they were in.
5. Analysis of different glucose-lowering medications is difficult as ACCORD patients were randomised to strategies, not to individual drugs.
a higher rate of events, they were even less likely to benefit from intensive glucose control than primary prevention patients. However, among those with no prior CVD, intensive treatment reduced the rate of events from $18 \%$ to $15.6 \%$, giving a significant absolute risk reduction of $2.4 \%$ (NNT 40), while among those with preexisting disease the absolute risk reduction was a non-significant $1.1 \%$. These results may suggest that tight glycaemic control influences the development of atherosclerosis but not its consequences.

## The ACCORD Study

The ACCORD (Action to Control Cardiovascular Risk in Diabetes) Study Group set out to address the advantages or otherwise of tight glycaemic and blood pressure control (assessed separately) in a range lower than that currently recommended, with $\mathrm{HbA}_{1 c}$ targeted below 6\% (ACCORD Study Group, 2008). The ACCORD study randomised over 10000 people with well-established diabetes (median duration 10 years) and either established CVD, or two risk factors for CVD. As in the ADVANCE trial, intensively treated participants had more visits to their health professional (every 1-2 months, with telephone contact between visits). Target $\mathrm{HbA}_{1 \mathrm{c}}$ was $<6.0 \%$ in the intensive group, $7-7.9 \%$ for the control group.

The glucose-lowering arm of the ACCORD study was halted in February 2008 (National Heart Lung and Blood Institute, 2008), due to a significant excess of all-cause mortality among intensively treated patients. Participants had spent between 3 and 5 years in the study, and the overall mortality was $1.41 \%$ in the intensive group and $1.14 \%$ among the standard treatment group. The absolute risk reduction was $2.7 \%$, relative risk increase was $22 \%(P=0.04)$. That is, as an apparent result of intensive blood glucose control, $22 \%$ more patients in the intensive arm died than those with more relaxed glucose targets and less medical attention.
Following the study being halted, sub-analysis of the ACCORD data is being undertaken, and

| Table 2. Comparison of mortality rates among patients who did and did not experience one or <br> more severe hypoglycaemic events during the ACCORD study. |  |  |
| :--- | :---: | :---: |
| Group | No hypoglycaemic events (\%) | Hypoglycaemic events (\%) |
| Overall mortality | 1.2 | 3.3 |
| Intensive group | 1.3 | 2.8 |
| Control group | 1.1 | 4.9 |

aims to answer a number of questions.

- Were certain groups of patients more or less likely to benefit or be harmed by intensive glycaemic control?
- Were any of the medications used responsible?
- Have we really reached the bottom in our efforts to remove the residual CV risk carried by people with type 2 diabetes?

It is important to be cautious when looking at sub-analyses as they can give a false sense of association. For example, if lower $\mathrm{HbA}_{1 \mathrm{c}}$ was found to be directly harmful in the ACCORD study, most glucose-lowering treatments would be implicated by association because they were being used more often in patients who are achieving lower $\mathrm{HbA}_{1 \mathrm{c}}$ levels.

Prior CV risk is not a major concern because all relevant risk factors were very uniform between the intensive and control groups in this large randomised study. However, twice as many participants in the intensive group as in the control group ( $28 \%$ versus $14 \%$ ) gained more than 10 kg of weight during the study, which might have led to a divergence in risk.

Hypoglycaemia in this study is a source of strong interest, especially in light of some evidence that severe hypoglycaemia is associated with adverse outcomes in other studies (Amiel et al, 2008; Shorr et al 1997; Campbell, 1985). Table 2 compares mortality rates among those who did and did not experience one or more severe hypoglycaemic events during the study.

These fascinating data tells us two things: Firstly, people experiencing severe hypoglycaemia were much more likely to die regardless of which treatment group they were in; and, secondly, among those who had experienced one or more serious event(s) post-randomisation, the intensive group fared significantly better. This seems to suggest that hypoglycaemia was not the mechanism by which intensively treated patients were carried to an early demise, but mortality in this group was in fact associated with some other causal factor.

What about social, educational and behavioural factors? In fact, insulin was used in $77 \%$ of intensively treated patients ( $55 \%$ of control), and bolus insulin in $40 \%$ (versus $20 \%$ of control). Self-monitoring of blood glucose was prescribed 3-8 times daily among the intensive group, including post-prandial monitoring. The
study authors do not indicate whether insulin self-adjustment was encouraged, but certainly the potential for erratic self-medication existed among the intensively treated patients.
Analysis of different glucose-lowering medications is difficult as patients were randomised to strategies, not to individual drugs. When duration of follow up on each drug was accounted for, there was a significant association with excess mortality only for patients on premixed and bolus insulins; a non-significant association existed for patients on glyburide, metformin, rosiglitazone and basal insulin; a non-significant improvement in survival was associated with use of pioglitazone; and a significant improvement was associated with the use of exenatide (although use of this drug occurred later in the study and was of generally short duration). Rosiglitazone use was associated with slightly lower mortality in intensively treated patients with pre-existing CVD and longer duration of diabetes.

## Discussion

What are the implications of these trails' outcomes for the primary care practitioner and the person with diabetes?

While intensive glucose-lowering does significantly reduce the risk of microvascular complications, and slows the progression of early complications, it does not significantly reduce CV events. On its own, this finding would continue to support intensive glucose-lowering for all patients, but the unexplained excess mortality among intensively treated patients in the ACCORD study suggests that an $\mathrm{HbA}_{1 \mathrm{c}}$ target greater than $6 \%$ is more appropriate. Selecting an intermediate $\mathrm{HbA}_{1 \mathrm{c}}$ target is difficult, and the results of the studies discussed above suggest that individual target-setting is more relevant than ever. We need to consider factors that include the relative burden of microand macrovascular risks, age and duration of diabetes. We face a growing population of younger people with type 2 diabetes who will live with the condition for even longer periods of time ( 30 years-). For these patients end-stage microvascular complications are a real longterm risk. In contrast, older patients who are unlikely to survive diabetes for more than 20 years, are much less likely to reach that stage of microvascular deterioration, and less stringent $\mathrm{HbA}_{1 \mathrm{c}}$ targets alongside aggressive CV risk
reduction will be more sustaining.
The latest NICE guidelines published in May 2008 (National Collaborating Centre for Chronic Conditions, 2008) suggest a phased approach, with lower $\mathrm{HbA}_{1 \mathrm{c}}$ targets around $6.5 \%$ in diabetes' early years when glucose control is achievable with diet and moderate doses of a single oral medication, moving to a less stringent $\mathrm{HbA}_{1 \mathrm{c}}$ target of around $7.5 \%$ as treatment becomes more complex. Primary care teams will likely consider this strategy more practical, more achievable and less stressful for all concerned than the 'as low as possible' approach with all the guilt that limited achievement usually entails.

Two other conclusions jump out from a close reading of these large studies. Firstly, $\mathrm{HbA}_{1 \mathrm{c}}$ levels tend to remain quite horizontal in these studies, in contrast to the inexorable rise throughout the UK Prospective Diabetes Study (UKPDS; 1998), demonstrating the effectiveness of regular review and stepwise intensification of glucose-lowering therapy. Secondly, the risk of a CV event is falling short of that predicted when these studies were being designed, which is an important marker of the success of multifactorial targeted risk reduction for CV events in diabetes from the time of diagnosis; success not only in design, but in execution.

As a GP, this author finds it undesirable to burden people with diabetes with a large number of drugs, but the evidence continues to support this line of action.

Type 2 diabetes continues to impose an excess risk of CVD aside from changes in CV risk factors - a state of affairs for which we currently have no solution.

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## Page points

1. The unexplained excess mortality among intensively treated patients in ACCORD study suggests that a $\mathrm{HbA}_{1 \mathrm{c}}$ target $>6 \%$ is more appropriate.
2. These new studies indicate that while intensive glucose-lowering does significantly reduce the risk of microvascular complications, it does not significantly reduce CVD events.
3. Selecting an intermediate $\mathrm{HbA}_{1 \mathrm{c}}$ target is difficult, and the results of these studies suggest that individual target-setting is more relevant than ever.
4. The latest NICE guidelines suggest a phased approach to $\mathrm{HbA}_{1 \mathrm{c}}$ targets. Primary care teams will likely consider this strategy more practical and achievable.
5. The $\mathrm{HbA}_{1 \mathrm{c}}$ levels in these studies tend to remain quite horizontal, demonstrating the effectiveness of regular review and stepwise intensification of glucose therapy.

[^0]:    Amiel SA et al (2008) Diabetic Medicine 25: 245-54
    ACCORD Study Group (2008) NEJM 358: 2545-59
    American Diabetes Association (2008) Press release: Intense Blood Glucose Control Yields no Significant Effect on CVD Reduction in VA Diabetes Trial. Available at: http://www.diabetes.org/for-media/pr-intense-blood-glucose-control-yields-no-significant-effect-on-cvd-reduction.jsp (accessed on 11.08.08)

    ADVANCE Collaborative Group (2007) Lancet 370: 892-40
    ADVANCE Collaborative Group (2008) NEJM 358: 2560-72
    Campbell IW (1985) Hormone and Metabolic Research Supplement Series 15: 105-11
    Kenny C (2008) Diabetes \& Primary Care 10: 66-68
    National Heart Lung and Blood Institute (2008) Press release: For Safety, NHLBI Changes Intensive Blood Sugar Treatment Strategy in Clinical Trial of Diabetes and Cardiovascular Disease. Available at: http://public. nhlbi.nih.gov/newsroom/home/GetPressRelease.aspx?id=2551 (accessed 11.08.08)

    National Collaborating Centre for Chronic Conditions (2008) Type 2 diabetes: national clinical guideline for management in primary and secondary care (update). Royal College of Physicians, London
    Shorr RI et al (1997) Archives of Internal Medicine 157: 1681-6
    UK Prospective Diabetes Study Group (1983) Diabetologia 24: 404-11
    UK Prospective Diabetes Study Group (1998) Lancet 352: 837-53

