Impact of data from recent clinical trials on strategies for treating patients with type 2 diabetes mellitus

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Abstract: Type 2 diabetes is associated with increased risk for the development of cardiovascular disease (CVD) secondary to hyperglycemia’s toxicity to blood vessels. The escalating incidence of CVD among patients with type 2 diabetes has prompted research into how lowering glycated hemoglobin ($HbA_1c$) may improve CVD-related morbidity and morality. Data from recent studies have shown that some patients with type 2 diabetes actually have increased mortality after achieving the lowest possible $HbA_1c$ using intensive antidiabetes treatment. Multiple factors, such as baseline $HbA_1c$, duration of diabetes, pancreatic β-cell decline, presence of overweight/obesity, and the pharmacologic durability of antidiabetes medications influence diabetes treatment plans and therapeutic results. Hypertension and dyslipidemia are common comorbidities in patients with type 2 diabetes, which impact the risk of CVD independently of glycemic control. Consideration of all of these risk factors provides the best option for reducing morbidity and mortality in patients with type 2 diabetes. Based on the results of recent trials, the appropriate use of current antidiabetes therapies can optimize glycemic control, but use of intensive glucose-lowering therapy will need to be tailored to individual patient needs and risks.

Keywords: type 2 diabetes, diabetes treatment, incretin-based therapies, glucose control, $HbA_1c$, cardiovascular disease

Introduction

Diabetes mellitus affects nearly 25 million people in the United States, with more than 90% diagnosed with type 2 diabetes.¹ Individuals with type 2 diabetes are at risk for a multitude of metabolic abnormalities that lead to microvascular and macrovascular complications, with cardiovascular disease (CVD) being the leading cause of mortality in these patients.² Despite advances in the diagnosis and treatment of CVD, mortality has increased in patients with type 2 diabetes at the same time as it has decreased in the general population.¹

Because of this alarming trend in patients with type 2 diabetes, reducing CV risk factors, including overweight/obesity, elevated blood pressure (BP), and dyslipidemia, is just as important as reducing hyperglycemia for maximizing outcomes in this patient population.²⁴⁻⁸ The importance of addressing these issues through individualized patient treatment strategies has been confirmed in a number of recent, large-scale clinical trials involving patients with type 2 diabetes.⁹⁻¹³

This paper will review data from recently conducted, large-scale clinical trials that evaluated the relationship between duration of disease, extent of glucose lowering, and cardiometabolic risk/CVD outcomes, and the treatment effects of more recently approved antidiabetes agents. The implications of these data on changes in current type 2 diabetes treatment practices will also be reviewed.
Optimum intensity of glycemic therapy in type 2 diabetes

Although epidemiologic studies indicate an association between elevated glycated hemoglobin (HbA$_{1c}$) and CVD in patients with type 2 diabetes, the effects of intensive glucose lowering on vascular outcomes remain unclear. Large-scale clinical trials enrolling patients with type 2 diabetes, including the Action to Control Cardiovascular Risk in Diabetes (ACCORD) study, the Action in Diabetes and Vascular Disease: Preterax and Diamicron MR Controlled Evaluation (ADVANCE), and the Veterans Administration Diabetes Trial (VADT), were designed to determine whether intensive antidiabetes therapy could reduce CVD events in this patient population. These studies are summarized in Table 1.

The results from these randomized clinical trials showed improved glycemic control (as measured by HbA$_{1c}$) with a significant difference demonstrated between the intensive antidiabetes therapy and the standard therapy groups. Blood pressure and serum lipid levels improved with appropriate administration of antihypertensive and dyslipidemia treatments with the antidiabetes therapies. While weight gain was noted in ACCORD and VADT in the intensive antidiabetes therapy group compared with the standard therapy group, there was weight loss in both treatment groups in ADVANCE, with the greater loss occurring in the standard therapy group (−1 kg) versus the intensive therapy group (−0.1 kg). Further, ACCORD and ADVANCE were secondary prevention trials for CVD in patients with type 2 diabetes and CVD and/or high risk for CVD, while the VADT study was a primary prevention trial for CVD in veterans with type 2 diabetes.

Data from ACCORD, ADVANCE, and VADT showed that reduction of CVD risk factors in patients with type 2 diabetes is not entirely dependent on the extent of glucose lowering. Instead, other factors, including disease duration and the presence of CVD comorbidities, have an influence on the morbidity and mortality of patients in this population. Intensive therapy in ACCORD was discontinued after a mean 3.5-year follow-up because of increased mortality in this treatment group.

Disease duration, comorbidities, and treatment outcomes in type 2 diabetes

Vascular complications and disease duration

Studies such as the Diabetes Control and Complications Trial (DCCT), the Wisconsin Epidemiologic Study of Diabetic Retinopathy (WESDR), and the United Kingdom Prospective Diabetes Study (UKPDS) have investigated the influence of disease duration and concomitant comorbidities on treatment outcomes.

The DCCT evaluated whether intensive antidiabetes treatment could decrease the frequency and severity of microvascular complications in patients with type 1 diabetes. Data showed that lowering HbA$_{1c}$ was associated with a reduced relative risk of microvascular complications, with the greatest reductions in patients with HbA$_{1c}$ > 9%. These findings are also important to patients with type 2 diabetes, as the pathophysiologic mechanisms driving disease progression are similar in type 1 and type 2 diabetes. There was no significant difference in macrovascular complications (CVD or mortality) between the standard and intensive therapy groups. However, a majority (93%) of the DCCT participants were subsequently followed from 1993–2005 for a mean follow-up of 17 years through the Epidemiology of Diabetes Interventions and Complications (EDIC) study. This study concluded that intensive diabetes therapy was independently associated with a significant decrease in the risk of CVD after the DCCT even though there was no significant difference in the HbA$_{1c}$ between the treatment groups in subsequent follow-up.

The WESDR examined the 25-year cumulative incidence and duration of macular edema (ME) (a commonly encountered microvascular complication in patients with type 1 and type 2 diabetes), and its association with various CVD risk factors, including hyperglycemia and BP. Data showed that elevated HbA$_{1c}$ and systolic BP were associated with an increased incidence of diabetic ME ($P < 0.004$ for both) in patients with type 1 and type 2 diabetes. As the duration of diabetes increased, the cumulative incidence of clinically significant ME and all-cause mortality also increased. Adjustment of data by patient age and gender showed that clinically significant ME was associated with increased CVD mortality in patients diagnosed with diabetes when they were aged ≥30 years. These data suggest that disease duration and the number of diabetes- and CVD-related comorbidities increase the morbidity and mortality of patients with type 2 diabetes.

Data from other clinical studies have shown that progression of type 2 diabetes and its related risk factors are favorably influenced by early initiation of treatment. In the UKPDS, newly diagnosed patients with type 2 diabetes (median baseline HbA$_{1c}$ 7.9% to 8.9%) were randomized to receive conventional glucose control (diet) or intensive glucose control (sulfonylurea [SFU], insulin, or metformin [MET]). Microvascular risk was reduced between 25%
to 29% in the intensive control group compared with the conventional control group during the intervention phase of the study and remained diminished throughout the 10-year post-trial phase despite a convergence of HbA1c in the two treatment groups.19 Moreover, although not significant during the interventional phase of the trial, patients in the SFU-insulin group experienced statistically and clinically relevant post-trial reductions in the risk for myocardial infarction (MI) (15%, \( P = 0.01 \)) and all-cause mortality (13%, \( P = 0.007 \)).

Patients with hypertension in the UKPDS were also randomized to stringent (angiotensin-converting enzyme inhibitor or \( \beta \)-blocker) or less-rigid BP control regimens (without these medications).20 During a 6- to 10-year interventional phase, mean BP was significantly lowered from baseline (146/81 mm Hg) to 143/79 mm Hg in the intensive control group compared with a slight increase to 152/82 mm Hg in the less-rigidly controlled group (\( P < 0.001 \) between treatment groups).20 These improvements in BP were associated with reduced risk of MI and microvascular disease.21

Unfortunately, the differences in CVD risk reduction between the two treatment groups were not sustained once the interventional trial ended.20 These data suggest that while early initiation of treating hyperglycemia and vascular complications/comorbidities is associated with improved morbidity and mortality in patients with type 2 diabetes, risk factor controls must be maintained to sustain long-term beneficial outcomes.

### Other comorbidities and type 2 diabetes

A number of studies (Look AHEAD [Action for Health in Diabetes] trial, Framingham Heart study, Diabetes Mellitus Insulin Glucose Infusion in Acute Myocardial Infarction [DIGAMI] studies, the Bypass Angioplasty Revascularization Investigation 2 Diabetes [BARI 2D] trial) have evaluated the impact of comorbidities, such as overweight/obesity and pre-existing CVD, on the morbidity and mortality of patients with type 2 diabetes.

In Look AHEAD, 5145 patients with a body mass index (BMI) > 25 kg/m\(^2\) were treated with either intensive lifestyle
intervention (ILI) involving increased physical activity and caloric restriction, or diabetes support and education (DSE) requiring attendance at support meetings.21 At the end of one year, ILI was associated with clinically significant weight loss, improved diabetes control, reduced CVD risk factors, and a reduction in the use of glucose-lowering medication compared with DSE.23 HbA1c decreased from 7.3% to 6.6% in the ILI group and from 7.3% to 7.2% in the DSE group ($P < 0.001$). High density lipoprotein-cholesterol (HDL-C) increased from baseline (44 mg/dL) in both groups after one year, but was significantly more improved in the ILI group (47 mg/dL) compared with the DSE group (45 mg/dL, $P < 0.001$).21

CVD is frequently present in overweight/obese patients with type 2 diabetes. Individuals with and without obesity and diabetes from the original and offspring cohorts of the Framingham Heart study were assessed for their lifetime risk of CVD.22 Over a 30-year period, the lifetime risk of CVD among normal-weight and obese females with diabetes was 54.8% and 78.8%, respectively.22 Among normal-weight and obese males with diabetes, the lifetime risk of CVD was 78.6% and 86.9%, respectively.22 These data showed that the lifetime risk of CVD is higher in patients with type 1 or type 2 diabetes and is further accentuated with increasing adiposity.

The DIGAMI 1 study established that initial intensive metabolic control with intravenous insulin followed by long-term subcutaneous insulin improved clinical outcomes in patients with type 2 diabetes and a prior MI.23 The DIGAMI 2 study was conducted to determine whether improvement was due to initial insulin-glucose infusion or to long-term subcutaneous insulin treatment.23 DIGAMI 2 compared three specific glucose-lowering regimens in patients with type 2 diabetes or elevated blood glucose and a suspected acute MI upon hospital admission: 1) a 24-hour insulin-glucose infusion followed by long-term subcutaneous insulin; 2) the same initial infusion treatment followed by standard glucose control; and 3) standard glucose control without the initial infusion treatment.24 Data demonstrated that although hypoglycemic events occurred most often in the first 24 hours after insulin treatment ($N = 111; 12\%$) compared with standard glucose control, hypoglycemia during time of admission was not associated with adverse CV outcomes.25 Only body weight (odds ratio [OR], 0.97; 95% confidence interval [CI]: 0.95 to 0.98; $P < 0.0001$) and diabetes duration (OR, 1.03; 95% CI: 1.01 to 1.05; $P = 0.0085$) were individually predictive of hypoglycemia and future morbidity and mortality.25

Upon discharge, patients enrolled in DIGAMI 2 were administered various antidiabetes agents, including insulin, MET, and SFUs for a median of 2.1 years. The results from post-hoc analyses demonstrated that insulin had a greater risk (updated, adjusted hazard ratio [HR], 1.73; 95% CI: 1.26 to 2.37; $P = 0.0007$) than MET (HR, 0.63; 95% CI: 0.42 to 0.95; $P = 0.03$) and SFU (HR, 0.81; 95% CI: 0.57 to 1.14; $P = 0.23$) for causing nonfatal MI and stroke in patients with type 2 diabetes.24 Additionally, further subanalysis of DIGAMI 2 data showed that insulin treatment after MI was associated with significantly increased weight gain (+2.3 kg; 95% CI: 1.5 to 3.2) and a 2.5-times greater incidence of reinfarction.26

Similarly, the BARI 2D trial evaluated whether revascularization and intensive antidiabetes therapy with either insulin-sensitization or insulin-provision could improve CV outcomes when compared with intensive medical therapy alone in patients with type 2 diabetes and stable coronary artery disease.27 After five years of treatment, survival rates did not differ significantly between revascularization (88.3%) and medical therapy (87.8%) groups or between the insulin sensitization (88.2%) and insulin provision (87.9%) groups. There was also no significant difference in reduction of CV events between the revascularization group (77.2%) and medical treatment group (75.9%) or between the insulin sensitization group (77.7%) and insulin provision group (75.4%).28 The results from these trials suggest that multiple factors beyond pre-existing conditions, including disease and comorbidity duration and intensity of selected therapy, interact to affect clinical outcomes in patients with type 2 diabetes.

**Multifactorial intervention**

Data from clinical studies have shown that treatment with multiple drug combinations can lower the risk of nonfatal CVD in some patients with type 2 diabetes. Steno-2 was conducted to evaluate the impact of this approach on CV risk factors and any-cause and CV-related mortality. Patients ($N = 160$) with type 2 diabetes and persistent microalbuminuria were treated with either intensive multifactorial intervention (stringent glycemic regulation, treatment with lipid- and BP-lowering agents, and aspirin) or conventional therapy for a mean of 7.8 years and followed-up for a mean of 5.5 years.12 Intensive multifactorial intervention resulted in significantly lower HbA1c compared with conventional therapy (7.9% vs 9.0%; $P < 0.01$), but these differences were no longer significant at the end of follow-up (7.7% vs 8.0%).12 After intervention, mean BP was reduced significantly with intensive (131/73 mm Hg) therapy compared with conventional (146/78 mm Hg) therapy, from 146/85 mm Hg and 149/86 mm Hg, respectively ($P < 0.01$ between treatment groups); significance was not maintained.
during the follow-up period. Fasting serum cholesterol was also significantly reduced with intensive (159 mg/dL) therapy compared with conventional (216 mg/dL) therapy, from 210 mg/dL and 233 mg/dL, respectively (P < 0.01 between treatment groups); again, significant improvements were not maintained after follow-up.12

Steno-2 data also showed that intensive multifactorial therapy was associated with sustained and beneficial effects on vascular complications, and any-cause, and CVD-related mortality. Nine patients in the intensive therapy group died from CVD-related causes compared with 19 in the conventional therapy group (P = 0.03).12 Diabetic nephropathy, progression of diabetic retinopathy, and autonomic neuropathy were significantly improved in the intensive therapy group compared with the conventional therapy group over the 13.3 years of observation (P < 0.01 for all).12 Although these data indicate that multifactorial care for patients with type 2 diabetes lowers CVD risk factors, morbidity, and mortality, the prompt and intensive implementation of antidiabetes treatments remains a formidable challenge in this patient population.

**Treatment strategies for type 2 diabetes**

Guidelines for the treatment of type 2 diabetes have been developed by various professional societies, including the American Association of Clinical Endocrinologists (AACE) and the American Diabetes Association (ADA). The guidelines, which are summarized in Table 2, provide target goals for several factors including glucose, BP, and lipids.2,3 Clinicians are encouraged to consider the individual patient needs and to encourage lifestyle changes. Results from clinical studies indicate that these targets are generally reasonable and obtainable but should be individualized and tailored to the needs and abilities of the patient with type 2 diabetes. Adjustments may be necessary for patients aged 65 years or older with and without comorbidities and for other individuals with mental and physical health challenges in order to avoid the potential hazards associated with tight glycemic control.

A list of selected antidiabetes pharmacotherapies with their advantages and disadvantages are presented in Table 3. Many of the available medications, including insulin, SFUs, thiazolidinediones (TZDs), and glinides, are associated with weight gain.8 MET, α-glucosidase inhibitors, and dipeptidyl peptidase-IV (DPP-IV) inhibitors are considered weight-neutral,9 while the glucagon-like peptide-1 (GLP-1) receptor agonists, amylin, and amylin analogs, are associated with weight loss.9 Insulin therapy may be useful in the undernourished patient to improve nutritional status and weight while avoiding hypoglycemia. Several reviews discuss the mechanisms of action and impact of these medications on other CVD comorbidities.8,29,30

The ADA and the European Association for the Study of Diabetes has developed a consensus algorithm to help guide initiation and adjustment of diabetes therapy using these agents (Figure 1).8 In addition to lifestyle improvements, this algorithm recognizes that clinicians have several antidiabetes medications available to help achieve glycemic targets. MET, SFU, and insulin are considered tier 1 core therapies, while tier 2 therapies include pioglitazone and a GLP-1 receptor agonist.4 Rosiglitazone is not recommended for use, and DPP-IV inhibitors are not listed. If tier 1 medications do not offer optimal benefits/glucose lowering, the use of additional agents, such as pioglitazone or a GLP-1 receptor agonist from tier 2, may be considered to provide glycemic

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**Table 2** Comparison of guidelines for the management of patients with type 2 diabetes

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>HbA1c</td>
<td>≤ 6.5%</td>
<td>&lt; 7%</td>
</tr>
<tr>
<td>Fasting glucose</td>
<td>Fasting plasma glucose &lt;110 mg/dL</td>
<td>Preprandial capillary plasma glucose, 70–130 mg/dL</td>
</tr>
<tr>
<td>Postprandial glucose</td>
<td>2-hr postprandial glucose &lt;140 mg/dL</td>
<td>Peak postprandial capillary plasma glucose &lt;180 mg/dL</td>
</tr>
<tr>
<td>BP</td>
<td>&lt; 130/80 mm Hg</td>
<td>&lt; 130/80 mm Hg</td>
</tr>
<tr>
<td>Lipids</td>
<td>LDL-C &lt;100 mg/dL (&lt;70 mg/dL for patients with diabetes and coronary artery disease)</td>
<td>LDL-C &lt;100 mg/dL</td>
</tr>
<tr>
<td></td>
<td>HDL-C &gt;40 mg/dL in men, &gt;50 mg/dL in women</td>
<td>HDL-C &gt;50 mg/dL</td>
</tr>
<tr>
<td></td>
<td>Triglycerides &lt;150 mg/dL</td>
<td>Triglycerides &lt;150 mg/dL</td>
</tr>
</tbody>
</table>

Notes: *In individuals with overt CVD, a lower LDL-C goal of <70 mg/dL (1.8 mmol/L), using high doses of a statin, is an option.**

Abbreviations: AACE, American Association of Clinical Endocrinologists; ACE, American College of Endocrinology; ADA, American Diabetes Association; BP, blood pressure; CVD, cardiovascular disease; HDL-C, high-density lipoprotein-cholesterol; LDL-C, low-density lipoprotein-cholesterol.
benefit. GLP-1 receptor agonists have the added benefit of promoting weight loss.8

The International Diabetes Center algorithm also includes goals and options for glycemic treatment.7 Beginning with complete lifestyle management, this algorithm recommends commencing treatment with MET and advancing to a two-drug therapy if target glycemic goals are not achieved within three months. The algorithm also includes guidance on the effect of antidiabetes therapy on hypoglycemia and weight (ie, among two drug therapies, only one promotes weight loss; the combination of MET and GLP-1 receptor agonist).7

A pathophysiology-based algorithm has been recently presented as an alternative to the ADA guidelines for the treatment of type 2 diabetes.6 This algorithm is based on targeting the pathophysiologic defects associated with diabetes, including impaired insulin secretion, increased lipolysis, decreased glucose uptake, and increased hepatic glucose production.6 Through lifestyle changes and triple therapy with a TZD, MET, and the GLP-1 receptor agonist exenatide, the regimen durably lowers hyperglycemia without inducing weight gain.6 Early commencement with this antidiabetes regimen should help to delay/prevent the progressive β-cell failure experienced by patients with type 2 diabetes.6

**Newer therapies for type 2 diabetes**

**Amylin agonist**

Patients with type 2 diabetes are deficient in the neuroendocrine hormone amylin, which is secreted by pancreatic β-cells (along with insulin) in response to nutrient intake.31 Amylin suppresses postprandial glucagon secretion and regulates gastric emptying and appetite.31 Treatment with pramlintide, a synthetic amylin agonist, as adjunctive therapy to insulin with or without oral antidiabetes agents, has been associated with improvements in glycemic control (up to −0.7% reduction in HbA1c), weight (up to −1.6 kg), and selected markers of CV risk (postprandial excursions of glucose, nitrotyrosine, and oxidized low-density lipoprotein-cholesterol [LDL-C]), without increased risk of hypoglycemia in patients with type 2 diabetes.32-34

**GLP-1 receptor agonists**

Incretins are gastrointestinal hormones that stimulate insulin release from pancreatic β-cells. The “incretin effect” describes the increased insulin secretion resulting from oral compared with intravenous glucose administration.35 It has been estimated that the incretin effect represents between 30% and 60% of the peripheral venous insulin response.35 As this represents a considerable proportion of postprandial insulin release, interest in developing incretins as diabetes treatment has increased over the past several years.

GLP-1 is a potent incretin hormone that is rapidly secreted by gut cells following a meal,36 although the enzyme DPP-IV rapidly metabolizes GLP-1 to an inactive fragment.37 Thus, the GLP-1 signaling pathway has been leveraged as a focus for diabetes research in two ways. First, GLP-1 receptor agonists have been developed that are resistant to DPP-IV

### Table 3 Effects on HbA1c: advantages and disadvantages of oral and parenteral antidiabetes agents

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfonylureas</td>
<td>Rapidly effective</td>
<td>Weight gain, hypoglycemia (especially with glyburide [dibenclamide in the EU] and chlorpropamide)</td>
</tr>
<tr>
<td>Metformin</td>
<td>Weight neutral</td>
<td>GI side effects, contraindicated in patients with renal insufficiency</td>
</tr>
<tr>
<td>TZDs</td>
<td>Improved lipid profile (pioglitazone), potential decrease in MI (pioglitazone)</td>
<td>Weight gain, fluid retention, CHF, bone fractures, expensive, potential increase in MI (rosiglitazone)</td>
</tr>
<tr>
<td>α-Glucosidase inhibitors</td>
<td>Weight neutral</td>
<td>Frequent GI side effects, TID dosing</td>
</tr>
<tr>
<td>Glinides (meglitinides)</td>
<td>Rapidly effective</td>
<td>Weight gain, TID dosing, hypoglycemia</td>
</tr>
<tr>
<td>DPP-IV inhibitors</td>
<td>Weight neutral</td>
<td>Risk of pancreatitis, renal failure</td>
</tr>
<tr>
<td>Insulin</td>
<td>No dose limit, rapidly effective, improved lipid profile</td>
<td>Weight gain, multiple daily injections, monitoring, hypoglycemia</td>
</tr>
<tr>
<td>GLP-1 receptor agonist</td>
<td>Weight loss</td>
<td>Frequent GI side effects, risk of pancreatitis, renal failure</td>
</tr>
<tr>
<td>Amylin/amylin analogue (pramlintide)</td>
<td>Weight loss</td>
<td>Frequent GI side effects, TID dosing, long-term safety not established</td>
</tr>
</tbody>
</table>

**Abbreviations:** CHF, congestive heart failure; DPP-IV, dipeptidyl peptidase-IV; EU, European Union; GI, gastrointestinal; GLP-1, glucagon-like peptide-1; MI, myocardial infarction; TZDs, thiazolidinediones.
Strategies for treating patients with type 2 diabetes

Exenatide, the synthetic form of exendin-4, which is 53% homologous to human GLP-1, binds the GLP-1 receptor agonist on pancreatic β-cells to potentiate insulin secretion. Exenatide has been approved by the US Food and Drug Administration (FDA) for use as an adjunctive therapy to improve glycemic control in patients with type 2 diabetes. Exenatide was shown to improve glucose levels and a number of CVD risk factors, including body weight, triglycerides (TG), total cholesterol, HDL-C, LDL-C, systolic BP (SBP) and diastolic BP in patients with type 2 diabetes treated for ≥3 years. In patients with type 2 diabetes previously treated with MET, the addition of exenatide improved hyperglycemic clamp-derived measures of β-cell function more than insulin glargine (P < 0.0001). In addition, in a retrospective database study, treatment with exenatide was associated with significant reduction in mean body weight (–2.7 kg; P < 0.001), BMI (–0.9 kg/m²; P < 0.001), abdominal girth (–2.9 cm; P < 0.001), total cholesterol (–7.4 mg/dL; P < 0.001), TG (–16.7 mg/dL; P < 0.001), and SBP (–2.6/–1.2 mm Hg; P < 0.03).

In pooled datasets from two studies, the addition of exenatide to patients treated with MET and an SFU for six months resulted in more patients reaching an HbA₁c ≤ 6.5% with significantly greater reductions in body weight (up to –3.7 kg; P < 0.0001) and SBP (–7.2 mm Hg; P < 0.005) than in patients who received insulin glargine or biphasic insulin aspart. In two other studies, exenatide produced greater reductions in SBP than either biphasic insulin aspart (–4.9 mm Hg vs –0.5 mm Hg; P < 0.0001) or placebo (–1.7 mm Hg vs +0.4 mm Hg; P < 0.0005).

A long-acting, once-weekly formulation of exenatide (which lowers glucose via the same mechanism as the

**Figure 1** ADA/EASD consensus guidelines treatment algorithm for patients with type 2 diabetes. Reinforce lifestyle interventions at every visit; check HbA₁c every three months until HbA₁c is <7% and then at least every six months. The interventions should be changed if HbA₁c is ≥7%. Copyright © 2009. Adapted with permission from Nathan DM, Buse JB, Davidson MB, et al. Medical management of hyperglycemia in type 2 diabetes: a consensus algorithm for the initiation and adjustment of therapy: a consensus statement of the American Diabetes Association and the European Association for the Study of Diabetes. Diabetes Care. 2009;32(1):193–203.

**Notes:** Sulfonylureas other than glyburide or chlorpropamide. Insufficient clinical use to be confident regarding safety.

**Abbreviations:** ADA, American Diabetes Association; CHF, congestive heart failure; EASD, European Association for the Study of Diabetes; GLP-1, glucagon-like peptide-1.
approved BID formulation) is currently under regulatory review by the US FDA. A 30-week, noninferiority trial in patients with type 2 diabetes receiving MET, a SFU, a TZD, or a combination of two of these agents, was conducted to compare exenatide administered twice daily to once weekly. Both treatment groups had reductions in HbA1c by week 6. Beginning at week 10 and continuing through the remainder of the trial, exenatide once weekly was associated with greater reductions in HbA1c than twice-daily dosing. After 30 weeks, the mean HbA1c reduction was −1.9% for exenatide once weekly versus −1.5% for twice daily (P = 0.002). Weight loss was similar (−3.7 kg) with both treatment regimens. Incidence of transient treatment-related nausea was significantly greater in the twice-daily group (34%; 50/148) compared with the once-weekly group (26.4%; 39/148; P < 0.05), while transient injection site pruritus was more commonly reported in the once-weekly group (17.6%; 26/148 vs 14.4%; 2/145). These events subsided over the duration of the study.

Liraglutide, an injectable GLP-1 receptor agonist with 97% homology to human GLP-1 and partial resistance to DPP-IV (through amino acid substitution and palmitate side chain addition), is being investigated in a once-daily formulation, and has recently been approved for use in the European Union. In the 26-week Liraglutide Effect and Action in Diabetes (LEAD)-2 MET study, patients with type 2 diabetes on a stable regimen of MET were treated with liraglutide, glimepiride, or placebo. After 12 weeks of therapy, liraglutide and glimepiride were associated with greater reductions in HbA1c (−0.7% to −1.0%) than placebo (−0.09%; P < 0.0001). Liraglutide reduced body weight up to −2.8 kg compared with a +1 kg weight gain with glimepiride (P < 0.0001). Liraglutide was associated with a greater incidence of nausea, vomiting, and diarrhea than either glimepiride or placebo, although the events were transient and subsided over the course of the study.

In a 52-week study, patients with type 2 diabetes receiving liraglutide monotherapy had significantly greater reductions from baseline HbA1c (−8.3%) than patients receiving glimepiride monotherapy (−1.1% vs −0.5%; P < 0.001). Unlike glimepiride, treatment with liraglutide was associated with weight loss (approximately −2.3 kg) while patients on glimepiride gained approximately 1 kg (P = 0.0001).

DPP-IV inhibitors
Sitagliptin, an oral antidiabetes agent, has been shown to inhibit plasma DPP-IV activity by −90% after two hours and by −80% after 24 hours post-dose. Sitagliptin monotherapy was associated with significant (P < 0.001) improvement in HbA1c (up to −0.9%) in a 24-week study of patients with a mean 4.4-year duration of type 2 diabetes.

Sitagliptin has also been studied in type 2 diabetes patients with inadequate glycemic control on MET alone. After 26 weeks of therapy, sitagliptin was well tolerated and provided significant improvement in HbA1c compared with placebo (−0.7% vs −0.02%; P < 0.001). In patients receiving inadequate glycemic control on pioglitazone, sitagliptin provided significant decreases (−0.9%) from baseline HbA1c (−8.1%) compared with placebo (−0.2%; P < 0.001).

Saxagliptin, another DPP-IV inhibitor, was recently approved for use in type 2 diabetes by the US FDA. In clinical studies, saxagliptin has been demonstrated to be effective in glucose-lowering in patients with type 2 diabetes either as monotherapy or in combination with other agents (eg, MET, SFU).

Implications of treatment decisions on the reduction of CVD risk
Based on data from multiple clinical trials, the current HbA1c goals for patients with type 2 diabetes appear appropriate. However, HbA1c values lower than 6.5% to 7% may not provide any clinical advantages in certain patient populations and may, in fact, increase mortality in patients with an already elevated risk for CVD. Although weight loss may be difficult for some patients with type 2 diabetes to achieve, it is associated with improvements in CVD- and diabetes-related risks. Additionally, longer duration of type 2 diabetes is associated with poorer clinical outcomes. Because patients with type 2 diabetes commonly present with varying levels of these risks, it is important to customize antidiabetes treatments within the framework of recommended treatment guidelines. Evidence continues to accumulate in support of newer antidiabetes agents, such as incretin-based therapies (GLP-1 receptor agonists and DPP-IV inhibitors), which improve hyperglycemia and other incretin-based therapies address an additional hormone deficiency present in patients with type 2 diabetes and expand the options for optimizing glucose control and management of the disease. However, undue delay in initiating insulin therapy with deteriorating glycemic control in order to try a variety of new noninsulin therapies should be avoided. Earlier initiation of insulin treatment should be considered to correct rising glycemia, particularly in patients with type 2 diabetes of more than 10 years duration.
Conclusions

Patients with type 2 diabetes have an elevated risk of CVD secondary to their hyperglycemia. This risk is compounded by the presence of common comorbidities including overweight/obesity, hypertension, and dyslipidemia. Addressing these risks simultaneously has beneficial effects on treatment outcomes. Some of the current antidiabetes pharmacotherapies, such as MET, α-glucosidase inhibitors, and incretin-based therapies have beneficial effects on hyperglycemia and other surrogate markers of CVD risk without increasing weight. SFUs, TZDs, glinides, and insulin control glycaemia and reduce CV risk factors but are associated with weight gain.

The duration of diabetes has an independent and negative impact on CVD risk and is related to the progression of morbidity and mortality. As evidenced by the results from recent, large-scale clinical trials, Hba1c target goals do not need to be drastically revised for patients with type 2 diabetes. Prompt initiation of antidiabetes treatment and individualized clinical judgment for each patient is paramount in treating hyperglycemia and its comorbidities. Earlier initiation of insulin therapy to optimize glycemic control should be considered in the management of type 2 diabetes.

Disclosures

Pasquale J Palumbo has no disclosures or conflicts. Jonathan M Wert has served as a consultant for educational services for Amylin Pharmaceuticals, Inc. and Eli Lilly and Company.

References


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