ABSTRACT

Lifestyle modifications in conjunction with antidiabetes medications can produce near-normal blood glucose concentrations in patients with type 2 diabetes mellitus (T2DM). Because these patients have increased cardiovascular morbidity and mortality, treatment strategies should also address the cardiovascular aspects of the disease, including blood pressure, lipids, and body weight. Since the prevalence of these abnormalities is increasingly secondary to poor diet and sedentary lifestyles and because most patients with T2DM are overweight/obese, clinicians are encouraged to help patients reduce body weight while correcting hyperglycemia by selecting treatment options that improve both parameters. The glucose-lowering properties of insulin and sulfonylureas are well known but they are also associated with weight gain. Thiazolidinediones are associated with weight gain as well as edema. However, this weight gain may be more peripheral than central, which may mitigate the risks associated with increased body fat. Metformin, the consensus first-line drug for the treatment of patients with T2DM, is weight neutral. Newer antidiabetes agents include incretin-based medications, such as the glucagon-like peptide–1 receptor agonists, which tend to decrease weight, and the dipeptidyl peptidase–4 inhibitors, which are weight neutral.

KEY POINTS

Control of cardiovascular risk factors is as important as glycemic control in patients with T2DM.

Intensive glucose control has shown mixed results in terms of correlation with improved cardiovascular risk factors.

Newer agents target the fundamental pathophysiologic defects of T2DM, with beneficial effects on weight and other cardiovascular risk factors.

T2DM, WEIGHT GAIN OR OBESITY, AND CV RISK: A CHALLENGING TRIAD

Despite therapeutic advances in the diagnosis and treatment of diabetes and CVD over the last decade, the estimated number of persons in the United States older than 35 years with self-reported diabetes (with T2DM accounting for 90% to 95% of diagnosed cases) and CVD has increased from 4.2 million in 1997 to 5.7 million in 2005. The CV risk for patients with T2DM who have not had a CV event such as a myocardial infarction (MI) is similar to that of individuals without diabetes who have had a prior MI. Patients with T2DM have nearly double the mortality of those without the disease. Adding to their risk, about 80% of patients with T2DM are overweight or obese, conditions associated with worsened insulin resistance and increased CV risk and disease burden. Even a modest weight gain (5 kg) may increase the risk of coronary heart disease (CHD) by 30%, while associated changes in lipids and BP can increase the risk by another 20%

It is as important to control CV risk factors as it is
to control glycemia in patients with T2DM, and both are difficult to achieve. Data from a recent nationwide Norwegian survey showed that only 13% of patients with T2DM achieved study-defined target levels; i.e., glycated hemoglobin (HbA1c) less than 7.5%, BP less than 140/85 mm Hg, and total cholesterol/high-density lipoprotein (HDL-C) ratio less than 4.0.10

**BENEFITS OF MANAGING GLYCEMIA, WEIGHT REDUCTION, AND CV RISK FACTORS**

Several large studies, many ongoing, are generating data on the relationships among glycemia, weight reduction, and CV risk. It is well established that individuals with T2DM need aggressive risk factor reduction (glucose control, blood pressure management, and treatment of dyslipidemia) to optimize outcomes. However, characterization of the benefits of various components of risk factor reduction, particularly over many years, is only now occurring.

Results from the United Kingdom Prospective Diabetes Studies (UKPDS) showed the benefits and risks of pharmacologic glycemic control—essentially monotherapy with insulin or a sulfonylurea—compared with conventional dietary therapy in reducing diabetic complications in patients with newly diagnosed T2DM. In UKPDS 33, both insulin and sulfonylureas (intensive treatment) reduced the risk of microvascular end points (retinopathy, nephropathy) in patients whose median HbA1c was lowered to 7.0% at 10 years of follow-up, compared with patients who reached an HbA1c of 7.9%. However, intensive glycemic control did not translate into a statistically significant reduction in macrovascular complications, including MI, stroke, CVD, and death. Additionally, patients assigned to insulin had greater weight gain (+4.0 kg) than did patients assigned to receive the sulfonylurea chlorpropamide (+2.6 kg) or glyburide (+1.7 kg) \( (P < .01) \).11

The UKPDS showed that intensive treatment with metformin reduced the risk of T2DM-related end points compared with conventional treatment (primarily diet alone) in overweight patients.12 Although there were fewer patients in the metformin-treated subset \( (n = 342) \) than in the conventional treatment cohort, a secondary analysis showed that metformin was associated with less weight gain and fewer hypoglycemic episodes than either insulin or sulfonylurea therapy.12 Since HbA1c levels in the treatment groups were equal, the additional benefits seen with metformin in overweight patients with T2DM were not based solely on glycemic control.

The Action to Control Cardiovascular Risk in Diabetes (ACCORD) trial involved 10,000 individuals with T2DM. The primary outcome measure was a composite of CV events. The intensively treated group was controlled to a target HbA1c of less than 6.0%, with most patients receiving insulin. The trial was terminated early because an increased risk of sudden death was observed.13 A similar study, Action in Diabetes and Vascular Disease: Preterax and Diamicron Modified Release Controlled Evaluation (ADVANCE), evaluated more than 11,000 patients with T2DM, starting with a sulfonylurea-based regimen. In this study, there was no reduction in macrovascular events, but there was a reduction in nephropathy in the intensively treated group.14 In both studies, hypoglycemia and weight gain were more frequent in intensively treated patients; and in ACCORD, there were more episodes of severe hypoglycemia in the intensive-treatment group.13,14

The Veterans Affairs Diabetes Trial (VADT) evaluated the effect of intensive glucose control on CVD in 1,791 patients \( (\text{mean age}, 60 \text{ years}) \) with poorly controlled T2DM \( (\text{average duration}, 11.5 \text{ years}) \). The primary end points included MI, stroke, new or worsening congestive heart failure (CHF), limb amputation, and invasive intervention for coronary or peripheral arterial disease. The hazard ratio for these end points in the intensive-treatment group was 0.88 \( (95\% \text{ confidence interval [CI]}, 0.74 \text{ to } 1.05) \).15,16 Specifically, the following beneficial effects were achieved:

- HbA1c reduced by \(-1.0\% \text{ to } -2.5\% \) in absolute units
- Systolic BP (SBP) reduced by \(-4 \text{ to } -7 \text{ mm Hg} \)
- Diastolic BP (DBP) reduced by \(-7 \text{ to } -8 \text{ mm Hg} \)
- Low-density lipoprotein cholesterol (LDL-C) reduced by \(-27 \text{ to } -28 \text{ mg/dL} \)
- Triglycerides reduced by \(-44 \text{ to } -50 \text{ mg/dL} \)
- HDL-C increased by \(4 \text{ to } 5 \text{ mg/dL} \)

Despite these benefits, body weight increased approximately 9 to 18 lb \((4 \text{ to } 8 \text{ kg}) \) during therapy.15

Since overweight and obesity are independent risk factors for CHD and CVD in patients with T2DM,17 weight management is an integral component in treatment. In the Action for Health in Diabetes (Look AHEAD) trial, an intensive exercise and weight-loss program resulted in clinically significant \((P < .001)\) weight loss at 1 year in patients who had T2DM and a body mass index (BMI) greater than 25 kg/m² \((> 27 \text{ kg/m²} \text{ if receiving insulin})\).18 When compared with patients who received less structured, infrequent support and minimal education about diabetes, participants in the intensive program showed more weight loss, improved glucose control, decreased CV events, and reduced medicine use. The Look AHEAD trial is currently evaluating whether these improvements will continue to result in lower CV risk.

**PATIENT ADHERENCE AND SATISFACTION**

It is often challenging for patients with T2DM to adhere to their treatment regimens. The Diabetes Attitude,
improvement in treatment satisfaction.20 Similarly, a
with injectable insulin. The 14.5% of patients who
A multicenter, randomized, clinical trial examined the
diabetes symptoms.21 The results from these studies show
mediated by lower self-efficacy perceptions and increased
BMI and poor adherence also had depression, which was
in adults with T2DM found that patients with higher
the interrelation of adherence, BMI, and depression
found that while 78% of patients with T2DM adhered
social barriers to self-care in patients with diabetes and
Wishes, and Needs (DAWN) study examined psychoso-
cial barriers to self-care in patients with diabetes and
found that while 78% of patients with T2DM adhered
to their medications, only 39% achieved complete suc-
cess in at least two-thirds of their self-care domains.19
A multicenter, randomized, clinical trial examined the
correlates of treatment satisfaction, including body
weight, on patients’ appraisal of treatment satisfaction
with injectable insulin. The 14.5% of patients who
experienced a reduction in BMI reported systematic
improvement in treatment satisfaction.20 Similarly, a
cross-sectionally designed study (n = 99) that analyzed
the interrelation of adherence, BMI, and depression
in adults with T2DM found that patients with higher
BMI and poor adherence also had depression, which
was mediated by lower self-efficacy perceptions and increased
diabetes symptoms.21 The results from these studies show
a clear relationship between adherence with treatment
regimens and achievement of HbA1c goals.22

**RECENT DEVELOPMENTS IN T2DM MANAGEMENT: STRATEGIES TO REDUCE CV RISK**

Because excess weight and obesity are prominent fea-
tures of T2DM, it is important to use an antidiabetes
agent that does not induce unnecessary weight gain
(particularly central weight gain, which is thought to
be most atherogenic).23 Metformin, considered the first-
line agent for treatment of T2DM, is generally weight
neutral with a low level of hypoglycemia.24,25 Sulfony-
lureas, insulin, and thiazolidinediones (TZDs) are all
associated with weight gain, although newer-analogue
insulins may cause less weight gain than older agents.
TZDs, especially pioglitazone, are associated with
improvements in long-term beta-cell function and CV
risk factors despite weight gain.26,27

The newer antidiabetes agents belong to the dipep-
tidyl peptidase-4 (DPP-4) inhibitor and the glucagon-
like peptide–1 (GLP-1) receptor agonist therapeutic
classes and have been shown to be either weight neu-
tral (DPP-4 inhibitors) or to cause weight loss (GLP-1
receptor agonists).28

**Figure 1** illustrates the physiologic role of GLP-1,29
which induces glucose-dependent insulin secretion
after food intake by binding to specific receptors on
pancreatic beta cells, suppresses postprandial gluca-
gon from pancreatic alpha cells, reduces postprandial
plasma glucose (PPG) concentrations by delaying gas-
tric emptying, and diminishes appetite.28 The dimin-
ished secretion of GLP-1 in T2DM30,31 has led to the
development of two different treatment approaches.28
Since GLP-1 is rapidly degraded by DPP-4, GLP-1
receptor agonists have been developed to resist DPP-4
inactivation while exhibiting many of the actions of
endogenous incretin hormones.28,29 DPP-4 inhibitors
function as incretin enhancers by protecting endog-
enous GLP-1 and glucose-dependent insulinotropic
peptide, another incretin, from enzymatic break-
down.31–33 Unlike the GLP-1 receptor agonists, which
are administered subcutaneously (SC), DPP-4 inhibi-
tors are administered orally.

**Obesity and the incretin effect**

A study in healthy subjects and patients with T2DM
demonstrated that glucose tolerance and obesity
independently impair the incretin effect, resulting in
impaired insulin secretion and glucagon suppression.34
Obesity is considered a subclinical inflammatory
condition that releases chemokines, leading to insulin
resistance. **Figure 2** illustrates the interaction be-
 tween obesity, inflammation, and insulin resistance.35

Two recent studies showed that surgically induced
weight loss enhances the physiologic “incretin effect.” In
one study, obese individuals with T2DM whose weight
loss was secondary to bariatric surgery combined with
caloric restriction showed improved insulin sensitivity,
improved carbohydrate metabolism, and elevated levels
of adiponectin and GLP-1, all of which may reduce the
incidence of T2DM.36 In the other study, bariatric sur-
gery in morbidly obese individuals with T2DM improved
insulin secretion and ameliorated insulin resistance.37

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**FIGURE 1.** Actions of glucagon-like–peptide–1 (GLP-1) in peripheral tissues. Most of the effects of GLP-1 are mediated by direct interaction with GLP-1 receptors on specific tissues. However, the actions of GLP-1 in liver, fat, and muscle most likely occur through indirect mechanisms.

DPP-4 inhibitors
DPP-4 inhibitors such as sitagliptin and saxagliptin inhibit the enzymatic activity of DPP-4 and increase endogenous concentrations of GLP-1. Sitagliptin has been compared with placebo as monotherapy and has been studied in combination with other therapies.

In an 18-week study, sitagliptin monotherapy, 100 and 200 mg QD, significantly reduced HbA1c compared with placebo (placebo-subtracted HbA1c reduction, −0.60% and −0.48%, respectively) in patients with T2DM. Sitagliptin also significantly decreased fasting plasma glucose (FPG) concentration relative to placebo. Twelve weeks of sitagliptin monotherapy at dosages of 5, 12.5, 25, and 50 mg BID led to significant (P < .001) reductions in HbA1c compared with placebo. Sitagliptin also produced significant reductions in FPG and mean daily glucose concentrations across the doses studied.

Similar results were reported in other 12-week studies: 50 mg BID and 100 mg QD sitagliptin monotherapy significantly (P < .05) reduced HbA1c −0.39% to −0.56% and FPG concentration −11.0 to −17.2 mg/dL compared with placebo; sitagliptin 100 mg QD compared with placebo produced a least-squares mean change from baseline HbA1c of −0.65% versus 0.41% (P < .001) and FPG of −22.5 versus 9.4 mg/dL (P < .001).

Sitagliptin also has been studied in combination with other therapies. After 24 weeks, sitagliptin combined with pioglitazone significantly reduced HbA1c by −0.70% and FPG by −17.7 mg/dL (P < .001 for both) compared with placebo. In another 24-week study, 100 mg sitagliptin QD significantly improved glycemic control and beta-cell function (P < .05 for both) in patients with T2DM who had inadequate glycemic control with glimepiride or glimepiride plus metformin.

In addition to significantly reducing HbA1c, sitagliptin 100 and 200 mg QD produced only small differences in body weight relative to placebo: least-squares mean change from baseline sitagliptin 100 mg was −0.7 kg (95% CI, −1.3 to −0.1) and for 200 mg was −0.6 kg (95% CI, −1.0 to −0.2); for placebo it was −0.2 kg (95% CI, −0.7 to 0.2). These findings were consistent with those from another 24-week monotherapy study where sitagliptin produced weight loss of up to −0.2 kg and a 30-week study of sitagliptin added to ongoing metformin therapy. In the latter study, both sitagliptin and placebo resulted in weight reductions of −0.5 kg.

The effects of sitagliptin on lipids and BP have been reported in clinical studies in patients with and without T2DM. In one study of patients with T2DM, the addition of sitagliptin to metformin increased total cholesterol (+8.1 mg/dL), LDL-C (+9.2 mg/dL), and HDL-C (+1.8 mg/dL) but lowered triglyceride (−14.5 mg/dL) after 18 weeks of treatment (24-week data). Data from a small (n = 19) study in nondiabetic patients with mild to moderate hypertension showed that sitagliptin produced small reductions (−2 to −3 mm Hg) in 24-hour ambulatory BP measurements.

Another DPP-4 inhibitor, saxagliptin, with efficacy similar to that described for sitagliptin, was recently approved by the US Food and Drug Administration (FDA) for treatment of T2DM.

GLP-1 receptor agonists
Many of the GLP-1 receptor agonists developed or under development have glucoregulatory effects similar to GLP-1 but are resistant to degradation by DPP-4. Exenatide, an exendin-4 receptor agonist, has compared favorably with sitagliptin and with insulin analogues. Long-acting (once-weekly and once-daily) GLP-1 receptor agonists are under development.

In a 2-week, head-to-head study in metformin-treated patients with T2DM, exenatide had a greater effect than sitagliptin in lowering PPG and was more potent in increasing insulin secretion and reducing postprandial glucagon secretion. In contrast to sitagliptin, exenatide slowed gastric emptying and reduced caloric intake.

In two studies of patients treated with exenatide, on a background of either metformin alone or metformin plus a sulfonylurea, patients who received metformin lost more weight (−1.6 to −2.8 kg; P ≤ .01) and experienced more significant decreases from baseline HbA1c (−0.4% to −0.8%; P < .002) at 30 weeks than did patients who received placebo. In a 16-week trial...
of exenatide in patients previously treated with a TZD with or without metformin, exenatide reduced HbA1c −0.98%, fasting blood glucose −1.69 mmol/L, and body weight −1.51 kg.62

When compared with insulin analogues, exenatide has been associated with weight loss (−3 kg) while the insulin analogues were associated with weight gain (+3 kg).53 After 26 weeks, body weight decreased −2.3 kg with exenatide and increased +1.8 kg with insulin glargine.54 Similar results were found in a crossover non-inferiority trial, where the least-squares mean difference in weight change was significantly (P < .001) different (2.2 kg) between the treatments.55 When exenatide was compared with insulin aspart in an open-label, non-inferiority trial, there was a between-group difference in weight of −5.4 kg after 52 weeks.32

Exenatide has also demonstrated these benefits in open-label extension studies. After 2 years, mean HbA1c reductions of −1.1% from baseline were sustained (P < .05), and weight loss of −4.7 kg was maintained (P < .001).66 After 82 weeks, similar HbA1c decreases (−1.1%) and weight loss (−4.4 kg) were exhibited.57 Even after 3 years, these benefits were maintained in patients who remained on the drug (HbA1c reduction from baseline, −1.0%; weight loss, −5.3 kg [P < .0001 for both]).58

Long-acting formulations of GLP-1 receptor agonists are in clinical development; two of these are once-weekly exenatide and once-daily liraglutide. Exenatide once weekly has the advantage of less frequent dosing and has elicited greater reductions in HbA1c than exenatide BID. After 15 weeks of once-weekly administration, the 0.8-mg formulation reduced HbA1c −1.4% and the 2-mg formulation reduced it −1.7% (P < .0001 for both compared with placebo). Body weight was lowered −3.8 kg (P < .05 compared with placebo) with the 2-mg formulation.59 Compared with exenatide BID, exenatide 2 mg once weekly showed greater reductions in HbA1c (−1.9% vs −1.5%; P = .0023) after 30 weeks of therapy.60 In a 1-year noncomparative trial, treatment with exenatide once weekly improved HbA1c (−2.0%) and weight (−4.1 kg), as well as BP and lipid profiles compared with baseline.61

Liraglutide, a once-daily human analogue GLP-1 receptor agonist, is under review by the FDA.18 In a 26-week study of patients with T2DM, liraglutide was associated with reductions in HbA1c (mean, −1.04%; P = 0.067 compared with insulin) and body weight (mean, −2.5 kg; P < .001 compared with insulin) at dosages of 0.6 to 1.8 mg/day SC. Liraglutide produced a decline in SBP from 0.6 to 3 mm Hg but was not associated with a decrease in DBP.62 In a 52-week study comparing liraglutide with glimepiride monotherapy, liraglutide 1.2 mg was associated with an HbA1c reduction of −0.84% (P = .0014) and the 1.8-mg dose with a reduction of −1.14% (P < .0001) compared with −0.51% for glimepiride. SBP decreased −0.7 mm Hg with glimepiride compared with −2.1 mm Hg for liraglutide 1.2 mg (P = .2912) and −3.6 mm Hg for liraglutide 1.8 mg (P < .0118). Mean DBP fell slightly but not significantly in all treatment groups.63 No effects on lipid parameters were reported in these two liraglutide studies.

The Liraglutide Effect and Action in Diabetes (LEAD-6) trial was undertaken to compare exenatide (10 μg BID SC) and liraglutide (1.8 mg/day SC) as add-on therapy to metformin, a sulfonylurea, or a combination of both in 464 patients with T2DM. After 26 weeks of treatment, liraglutide was associated with a significant reduction in HbA1c of −1.12%, compared with −0.79% with exenatide (P < .0001). Patients treated with liraglutide lost −3.2 kg while those on exenatide lost −2.9 kg. Among patients previously treated with metformin alone, there was a 1-kg difference in favor of liraglutide (P = NS).64

### Safety profile

All of the drugs discussed have potential adverse effects. Metformin continues to have a black box warning for lactic acidosis.65 Sulfonylureas and insulin can cause hypoglycemia. TZDs can cause fluid retention and, in rare cases, CHF (for which these drugs also carry a black box warning).66,67 TZDs also increase the risk of distal fracture.66,67 The most common side effects of exenatide are gastrointestinal, but there have been reported cases of pancreatitis, some of which have been fatal.68,69 It has been difficult to prove whether exenatide increases the risk of pancreatitis, as patients with T2DM are already at an increased (three- to fourfold) risk for this condition compared with persons who do not have T2DM.69 Exenatide should not be used in patients with severe renal impairment or end-stage renal disease; it should be used with caution in patients who have undergone renal transplantation and in patients with moderate renal impairment.

The prescribing information for sitagliptin includes pancreatitis among the adverse reactions identified during the drug’s postapproval use.70 As with exenatide, it is not fully known whether a true association exists between the agent and pancreatitis. However, since pancreatitis can occur in this patient population, it is recommended that abdominal pain be fully evaluated to rule out pancreatitis. Continued postmarketing surveillance is important for all of these agents.

### The role of guidelines

The American Association of Clinical Endocrinologists (AACE),26 the American Diabetes Association (ADA),71 and the ADA in conjunction with the European Association for the Study of Diabetes (EASD)24 have recently revised their recommendations for the
management of patients with diabetes. The guidelines are unanimous in setting a glycemic goal (HbA1c < 7.0% for the ADA, HbA1c ≤ 6.5% for the AACE) and advocating individualized care for a treatment goal of HbA1c lower than 6.0% in patients who stand to benefit from near euglycemia without inducing severe hypoglycemia.24

CVD is the major cause of morbidity and mortality associated with T2DM and is a source of increasing concern. Accordingly, special consideration should be given to patients with coexisting CV risk factors, including hypertension and dyslipidemia. The ADA and the EASD advocate lifestyle modification to decrease body weight and the concurrent initiation of metformin as first-line therapy.24 If that strategy is insufficient, then two tiers of treatment guide the choice of next steps:24,25

- Tier 1, in addition to metformin, includes the sulfonylureas and insulin. Although these are excellent glucose-lowering drugs, they are associated with weight gain, hypoglycemia, and no improvement in BP or lipid levels. They are relatively low in cost and have been used for many years. Their main drawback is evidence that despite their use, beta-cell failure continues unabated over time.

- Tier 2 treatments include pioglitazone and the GLP-1 receptor agonist exenatide. Consideration may be given to the use of pioglitazone or exenatide when hypoglycemia is of concern, with exenatide being preferred when weight loss is a major objective and HbA1c is close to target (< 8.0%). Additionally, both the TZDs and exenatide probably help slow the rate of beta-cell failure, particularly if they are used early in the course of the disease.27 The AACE recommends different pharmacologic approaches based on HbA1c at diagnosis.26

The American Heart Association and the ADA have issued a joint scientific statement on the primary prevention of CVD in patients with diabetes. They advocate lifestyle management of body weight, nutrition, and physical activity. In addition, they stress the need for attention to BP, lipid levels, and smoking status, and the use of antiplatelet agents in patients at increased CV risk (> 40 years of age and a family history of CVD, hypertension, smoking, dyslipidemia, or albuminuria).

CONCLUSION

T2DM, weight gain/obesity, and CV risk present a continuing challenge to patients and clinicians. Antidiabetes agents have varying degrees of evidence to support their effects on HbA1c, body weight, BP, and lipid levels. A better understanding of the pathophysiology of T2DM has led to the development of newer antidiabetes agents that target the fundamental defects of the disease. Evidence continues to accumulate for the improved benefits of glycemic control and weight loss in T2DM with GLP-1 receptor agonists such as exenatide currently having robust data in terms of beneficial effects on weight and CV risk factors. As clinicians continue to incorporate this knowledge into their practice patterns, patient adherence and clinical outcomes are expected to improve. Newer agents, such as incretin-based therapies, address T2DM as well as other factors that increase cardiometabolic risk through their effects not only on glycemic control but on body weight, BP, and lipids.

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