Four-Year Analysis of Cardiovascular Disease Risk Factors, Depression Symptoms, and Antidepressant Medicine Use in the Look AHEAD (Action for Health in Diabetes) Clinical Trial of Weight Loss in Diabetes

OBJECTIVE—To study the association of depressive symptoms or antidepressant medicine (ADM) use with subsequent cardiovascular disease (CVD) risk factor status in the Look AHEAD (Action for Health in Diabetes) trial of weight loss in type 2 diabetes.

RESEARCH DESIGN AND METHODS—Participants (n = 5,145; age [mean ± SD] 58.7 ± 6.8 years; BMI 35.8 ± 8.5 kg/m²) in two study arms (intensive lifestyle [ILI], diabetes support and education [DSE]) completed the Beck Depression Inventory (BDI), reported ADM use, and were assessed for CVD risk factors at baseline and annually for 4 years. Risk factor–positive status was defined as current smoking, obesity, HbA1c ≥7.0% or insulin use, and blood pressure or cholesterol not at levels recommended by expert consensus panel or medicine to achieve recommended levels. Generalized estimating equations assessed within-study arm relationships of elevated BDI score (≥11) or ADM use with subsequent year CVD risk status, controlled for demographic variables, CVD history, diabetes duration, and prior CVD risk status.

RESULTS—Prior year elevated BDI was associated with subsequent CVD risk factor–positive status for the DSE arm (A1C [odds ratio 1.30 (95% CI 1.09–1.50]), total cholesterol [0.80 (0.65–1.00)], i.e., protective from high total cholesterol) and the ILL arm (HDL [1.40 (1.12–1.75)], triglyceride [1.28 (1.00–1.64)]). Prior year ADM use predicted subsequent elevated CVD risk status for the DSE arm (HDL [1.24 (1.03–1.50)], total cholesterol [1.28 (1.05–1.57)], current smoking [1.73 (1.04–2.88)] and for the ILL arm (A1C [1.25 (1.08–1.46)], HDL [1.32 (1.11–1.58)], triglycerides [1.75 (1.43–2.14)], systolic blood pressure [1.39 (1.11–1.74)], and obesity [1.46 (1.22–1.81)].

CONCLUSIONS—Aggressive monitoring of CVD risk in diabetic patients with depressive symptoms or who are treated with ADM may be warranted.

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Elevated rates of depression among people with diabetes (1) may partly account for their higher rates of cardiovascular disease (CVD) morbidity and mortality (2). Depression is associated with adverse CVD outcomes (3,4), likely via behavioral mechanisms (e.g., effects of cigarette smoking, and sedentary lifestyle and poor diet leading to obesity) and physiological mechanisms (e.g., effects of elevated blood glucose, blood pressure, and lipid levels, as well as dysregulation of the hypothalamic-pituitary-adrenal axis). Behavioral mechanisms may activate physiological mechanisms.

Literature suggests that antidepressant medicines (ADMs) may also affect CVD risk factors and outcomes either negatively or positively. Some ADMs, such as tricyclic antidepressants (TCAs), may increase the risk of myocardial infarction (5). Findings on the association between other widely used ADMs, such as selective serotonin reuptake inhibitors (SSRIs) and serotonin–noradrenaline (norepinephrine) reuptake inhibitors (SNRIs), and cardiovascular outcomes are mixed. Some have reported positive outcomes in patients with established CVD (6). Others have reported no such benefits (7–9).
In the Diabetes Prevention Program (DPP), ADM use was associated with increased risk of developing type 2 diabetes, raising the possibility that ADM use could lead to negative health outcomes in people with diabetes or at risk for the disease. DPP participants in the placebo and intensive lifestyle intervention (ILI) arms of the study were two to three times more likely to develop diabetes during the course of the study if they were taking ADMS than if they were not (10,11).

In 2010 we reported that among participants in the Look AHEAD (Action for Health in Diabetes) clinical trial, depression symptoms and ADM use on entry to the study were each independently associated with a wide range of CVD risk factors (12). Here we assessed the temporal dynamics of elevated depression symptoms and ADM use with selected CVD risk factors during the first 4 years of Look AHEAD. Our specific aim was to determine whether elevated depression symptoms or ADM use were independently associated with subsequent elevated CVD risk factors over trial years 1–4, controlled for baseline characteristics of age, sex, race/ethnicity, education, history of CVD, and diabetes duration, as well as CVD risk factor status in the prior year.

**RESEARCH DESIGN AND METHODS**—Look AHEAD is a randomized clinical trial of 5,145 overweight or obese individuals with type 2 diabetes designed to assess the long-term effect (up to 14 years) of a comprehensive behavioral weight loss intervention on cardiovascular and other health outcomes. Participants were randomized to an ILI or to a diabetes support and education (DSE) treatment arm. The ILI included goals for diet modification (1200–1800 kcal/day based on initial weight and physical activity (175 min of moderate physical activity per week), designed to induce at least 7% weight loss at year 1 and to maintain weight loss in subsequent years. ILI participants were seen weekly for the first 6 months and 3 times per month for the next 6 months. During years 2–4, participants were seen individually at least once a month and contacted another time each month by telephone or e-mail. DSE participants were invited to three group sessions each year. Sessions followed a standard protocol and covered diet, exercise, and social support, without addressing behavioral strategies.

Their personal physicians provided medical care for all participants. These physicians made changes in medications, with the exception of changes in diabetes medication made by Look AHEAD physicians when an ILI participant was losing substantial weight. Participants are being monitored at 16 clinical centers in the U.S.

Inclusion criteria for entry to the study were 1) age 45–76 years; 2) BMI ≥25 kg/m2 (27 kg/m2 if currently taking insulin because thinner individuals taking insulin may be less responsive to weight loss); and 3) glycosylated hemoglobin (HbA1c) <11%, systolic blood pressure (SBP) <160 mmHg, diastolic blood pressure (DBP) <100 mmHg, and triglyceride (TG) <600 mg/dL. Exclusion criteria were 1) underlying diseases or conditions likely to affect the safety of the interventions or factors that might limit adherence to the interventions or affect conduct of the trial, including hospitalization for depression in the past 6 months; suicidal ideation; current diagnosis of schizophrenia, other psychotic disorders, or bipolar disorder; or self-report of alcohol or substance abuse within the past 12 months; or 2) other medical, psychiatric, or behavioral limitations (e.g., difficulty completing the 2-week run-in period during which participants were required to record food eaten) that in the judgment of the principal investigator might interfere with study participation or the ability to follow the protocol. ADM use was not a criterion for exclusion, nor was having depression symptoms that did not require hospitalization in the prior 6 months or that did not involve suicidal ideation. Full details of the Look AHEAD design and methods are reported elsewhere (13); however, measures relevant to this report are briefly described below.

### Assessments

The following assessments were completed annually:

- **Serum measurements.** The Central Biochemistry Laboratory (Northwest Lipid Research Laboratories, University of Washington, Seattle, WA) conducted standardized analyses of HbA1c, fasting serum glucose, total serum cholesterol (TC), LDL cholesterol, HDL cholesterol, and TG in frozen samples (14).

- **Weight, blood pressure, and smoking.** Weight and height were measured in duplicate using a digital scale and stadiometer. BP was measured in duplicate with an automated device using standardized quality controlled protocols. Participants reported their smoking status (current, former, never).

- **Depression symptoms and ADM use.** Participants brought all prescription medicines to their assessment visits. Study staff recorded the name of each medicine; dosages were not recorded. At each visit, participants completed the Beck Depression Inventory (BDI) (15), a self-report scale with reliable psychometric characteristics across a broad spectrum of clinical and nonclinical populations. The BDI lists 21 symptoms, with responses scored from 0 to 3 in ascending symptom severity, and total scores ranging from 0 to 63, with higher scores indicating more symptom burden. Elevated depression symptoms were defined by a BDI score ≥11, a value used in the earlier Look AHEAD report and in the DPP reports (10–12).

- **CVD risk factor classification.** The primary objective of this report was to assess the association between elevated depression symptoms or ADM use and subsequent CVD risk factor–positive status. Each of nine elements of five CVD risk factors was dichotomized into risk–positive and not. Risk factor–positive status was defined as current smoking, BMI ≥30 kg/m2, HbA1c >7.0%, SBP >130 mmHg, DBP >80 mmHg, LDL ≥100 mg/dL, HDL ≤40 mg/dL, TC ≥200 mg/dL, and TG ≥150 mg/dL, as recommended by the American Diabetes Association (16) or the Expert Panel on Detection, Evaluation, and Treatment of High Blood and Cholesterol in Adults (17), or taking medicine to achieve these targets (18,19). Insulin was the only glucose-lowering agent considered as an indicator of CVD risk–positive status because almost all participants were taking some glucose-lowering medicine. Insulin use and A1C are commonly used composite measures of diabetes control. All antihypertensive and all lipid-lowering agents were considered as “at risk” indicators (the list of medicines included in this determination and the ADMS the participants took is presented in the Supplementary Data).

- **Outcomes.** We assessed the association that positive status for each CVD risk factor had with elevated BDI scores and ADM use in the prior year, controlled for CVD risk factor status of interest in the prior year, participant characteristics of age, sex, race, education, history of cardiovascular disease, and duration of diabetes, and year of follow-up.
Statistical analysis
Analyses included all randomized participants according to intervention assignment. First-order Markov models were used to parameterize inter-subject longitudinal correlations and were fitted using generalized estimating equations (20); higher order models were deemed unnecessary based on Wald tests. Because relationships differed between intervention treatment arms based on tests of interaction, odds ratios (ORs) with 95% CIs are reported separately for DSE and ILI participants. We also conducted a series of ancillary analyses.

RESULTS—Table 1 reports the characteristics of the study population by intervention assignment. Earlier, we reported that at baseline, 16.5% of participants were taking ADM and 14.7% had elevated depression symptom scores (BDI $\geq 11$), indicating likely mild to moderate depression, and 26.8% had elevated depression symptom scores or were using ADM (12). However, 85.3% had BDI scores $< 11$ (median 8; 25th–75th percentile, 6–9), reflecting that many individuals with severe depression symptoms resulting in hospitalization or inability to successfully complete the run-in period were excluded from Look AHEAD participation (12).

Elevated depression symptoms, ADM use, and CVD risk factor–positive status
Table 2 reports the proportions of DSE and ILI participants with elevated depression symptom scores, using ADM, and with elevated CVD risk factors at baseline and at annual assessments during the first 4 years of the study. Of the participants taking ADMs, 73% took SSRIs, SNRIs, or serotonin modulators, 28% took norepinephrine-dopamine reuptake inhibitors, and 23% took TCA or tetracyclic agents at some point during the 4-year follow-up. Thirteen percent of participants not using ADMs at baseline took them at some point during the 4-year follow-up. Twenty-one percent of those using ADMs at baseline stopped taking them during follow-up. Seventeen percent of participants with a BDI score $< 11$ at baseline had an elevated BDI score at some time during the 4-year follow-up. Of those with elevated BDI scores at baseline, 81% had a score below the cutoff of 11 at some point during follow-up. BDI scores were elevated in 27–29%, and 26–28% were taking ADMs at some point in the study. The proportion of participants with elevated BDI scores who were taking ADMs was 5.7, 7.2, 8.6, and 9.5% in years 1, 2, 3, and 4, respectively, in the DSE arm and 6.2, 7.9, 9.1, and 9.8%, respectively, in the ILI arm.

The proportion of participants with an elevated A1C level or taking insulin, as well as the proportion with a BMI $\geq 30$ kg/m$^2$, differed substantially between intervention groups. The proportion of DSE and ILI participants with an elevated A1C level or taking insulin at some point in the study was 75.6 and 67.4%, respectively. The proportion of DSE and ILI participants with a BMI $\geq 30$ kg/m$^2$ was 90.8 and 86.9%, respectively.

| Variable                     | Total $n = 5,145$ | DES arm $n = 2,575$ | ILI arm $n = 2,570$ |
|------------------------------|-------------------|---------------------|---------------------|
| Age (years)                  | 58.8 ± 6.9        | 58.6 ± 6.8          |
| Female sex                   | 1,537 (59.7)      | 1,526 (59.4)        |
| Race                         |                   |                     |
| African American/black (not Hispanic) | 803    | 404 (15.7)          | 399 (15.6)          |
| Native American*             | 258               | 128 (5.0)           | 130 (5.1)           |
| White                        | 3,253             | 1,628 (63.4)        | 1,618 (63.1)        |
| Hispanic                     | 681               | 338 (13.2)          | 339 (13.2)          |
| Other†                       | 150               | 50 (1.9)            | 48 (1.9)            |
| Education                    |                   |                     |
| <13 years                    | 1,024             | 515 (20.5)          | 509 (20.2)          |
| 13–16 years                  | 1,915             | 968 (38.6)          | 947 (37.5)          |
| >16 years                    | 2,094             | 1,027 (40.9)        | 1,067 (42.3)        |

Data are expressed as mean ± SD or n (%). *Includes American Indian and Alaskan native. †Includes Asian, Pacific Islander, mixed, and missing.

| Measure                | Baseline % | 1 % | 2 % | 3 % | 4 % | Any time % |
|------------------------|------------|-----|-----|-----|-----|------------|
| BDI $\geq 11$          | DSE 12.9   | 11.4 | 12.8 | 12.1 | 12.3 | 28.9       |
|                        | ILI 14.9   | 9.5  | 11.7 | 11.5 | 12.3 | 27.5       |
| ADM use                | DSE 15.3   | 17.1 | 16.8 | 19.0 | 19.3 | 26.2       |
|                        | ILI 17.6   | 17.5 | 18.6 | 19.9 | 20.1 | 27.8       |
| CVD risk factor positive |           |     |     |     |     |            |
| A1C >7.0% or insulin   | DSE 56.9   | 51.8 | 52.5 | 52.1 | 54.5 | 75.6       |
|                        | ILI 55.2   | 31.1 | 39.4 | 42.2 | 46.3 | 67.4       |
| LDL ≥100 mg/dl or medicine | DSE 80.0  | 88.5 | 90.1 | 89.3 | 89.9 | 96.3       |
|                        | ILI 88.5   | 87.3 | 88.0 | 87.2 | 89.6 | 95.5       |
| HDL ≤40 mg/dl or medicine | DSE 71.1  | 75.9 | 79.3 | 81.4 | 83.3 | 93.3       |
|                        | ILI 70.7   | 70.4 | 74.4 | 78.0 | 79.6 | 90.8       |
| TC ≥200 mg/dl or medicine | DSE 73.9  | 75.4 | 80.1 | 80.4 | 82.0 | 89.4       |
|                        | ILI 74.6   | 73.2 | 77.4 | 78.0 | 80.7 | 88.3       |
| TG ≥150 mg/dl or medicine | DSE 75.0  | 77.7 | 80.3 | 82.6 | 83.7 | 90.9       |
|                        | ILI 75.6   | 72.8 | 77.7 | 80.8 | 82.2 | 90.6       |
| SBP ≥130 mmHg or medicine | DSE 77.9  | 78.4 | 80.7 | 81.7 | 83.4 | 90.3       |
|                        | ILI 78.9   | 76.4 | 78.9 | 80.5 | 83.0 | 88.1       |
| DBP ≥80 mmHg or medicine | DSE 84.4   | 84.4 | 83.4 | 87.1 | 87.7 | 94.7       |
|                        | ILI 84.4   | 81.2 | 82.8 | 85.1 | 87.1 | 92.3       |
| Current smoking         | DSE 4.3    | 4.0  | 4.2  | 3.6  | 3.6  | 5.9        |
|                        | ILI 4.6    | 4.7  | 4.0  | 4.0  | 3.9  | 6.3        |
| BMI ≥30 kg/m²           | DSE 85.9   | 84.0 | 83.3 | 82.8 | 82.7 | 90.8       |
|                        | ILI 84.3   | 63.2 | 67.8 | 71.4 | 73.2 | 86.9       |
Associations between CVD risk factor status and prior year indicators of depression

Table 3 summarizes the ORs linking current CVD risk factor–positive status with indicators of depression at the prior annual examination for each of the nine risk factors in DSE and ILI participants, controlled for the CVD risk factor of interest status at the prior examination. ORs for CVD risk factor–positive status include those who were risk factor–positive the previous year and remained this way as well as those who were risk factor–negative the previous year and became risk factor–positive. In the DSE arm, only two associations with elevated BDI reached nominal statistical significance: the odds (95% CI) of elevated A1C/insulin use were increased (1.03 [1.09–1.56]) and the odds of elevated TC/medicine use were decreased if BDI was elevated in the prior year. Two associations with elevated BDI also reached statistical significance in the ILI arm: the odds of low HDL/medicine use (1.40 [1.12–1.75]) and elevated TC/medicine use (1.28 [1.01–1.64]) were increased if BDI was elevated in the prior year.

In the DSE cohort, ADM use in the prior year was associated with more prevalent low HDL/medicine use (1.20 [1.03–1.50]), elevated TC/medicine use (1.29 [1.05–1.57]), and current smoking (1.70 [1.04–2.88]). In the ILI arm, ADM use in the prior year was associated with more prevalent elevated A1C/insulin use (1.25 [1.08–1.46]), low HDL/medicine use (1.33 [1.11–1.58]), elevated TC/medicine use (1.75 [1.43–2.14]), elevated SBP/medicine use (1.39 [1.11–1.74]), and BMI ≥30 kg/m² (1.47 [1.22–1.76]).

We also conducted a series of ancillary analyses that confirmed the robustness of our original models (results not shown): First, repeating the analyses using continuous BDI scores rather than a dichotomous classification did not change the direction or statistical significance of any relationships with CVD risk factors.

Second, when we re-estimated the models with log-transformed BDI scores, we found similar relationships with almost identical ORs and 95% CIs.

Third, when we repeated the analyses excluding participants who took TCA or tetracyclic ADMs, the results were nearly identical to analyses involving individuals taking any ADM.

Fourth, we repeated the analyses controlled for weight change during the study. The pattern of associations with ADM use was similar to the pattern in our primary analysis. Six of eight associations (all but the association with smoking in the DSE arm and low HDL/medicine in the ILI arm) that were significant in the primary analysis were also significant in the analysis controlled for weight change.

In the ancillary analysis, only one of four associations with elevated BMI (the association with A1C >7.0% or insulin in the DSE arm) that had been significant in the primary analysis was significant in the analysis controlled for weight change. Tests for interaction between age and ADM and BMI yielded nonsignificant relationships (P > 0.05).

**Conclusions**—We found that depression markers (elevated depression symptom scores or ADM use) during the prior year were associated with current elevated CVD risk factors in both Look AHEAD intervention arms when prior risk factor status and other covariates were controlled. These significant associations were most common for ADM use in the ILI treatment arm, but the analyses reported here were not designed to explicate treatment arm differences in these associations. Thus, the available data do not allow us to determine the reasons for these findings.

We assessed CVD risk factors in the five domains of glycemia, lipids, blood pressure, smoking, and BMI, all of which are well-documented risk factors for CVD morbidity and mortality (21-25). We found that at least one indicator from each of these domains was increased in the presence of elevated depression symptoms or ADM use. Overall, there were more significant associations of CVD risk factors with ADM use than with depression symptoms.

We found little evidence for a temporal relationship between elevated depression symptoms and subsequent increases in positive CVD risk factor status. Others have reported that depression is associated with physiological abnormalities that could contribute to adverse cardiovascular outcomes, including abnormalities that are likely associated with depression per se, because they are observed in depressed patients who do not have CVD. These abnormalities include increased inflammatory markers, endothelial dysfunction, abnormal platelet activation, elevated catecholamine levels,

Table 3—ORs for CVD risk factor–positive status with ADM use or BDI ≥11 the preceding year, controlled for risk factor status in the prior year

| CVD risk factor | ADM use OR (95% CI) | P value | BDI ≥11 OR (95% CI) | P value |
|-----------------|---------------------|---------|--------------------|---------|
| **DSE arm**     |                     |         |                    |         |
| A1C >7.0% or insulin | 1.13 (0.96–1.32) | 0.14    | 1.30 (1.09–1.56) | 0.04    |
| LDL ≥100 mg/dL or medicine | 1.24 (0.96–1.60) | 0.10    | 0.86 (0.67–1.11) | 0.24    |
| HDL ≤40 mg/dL or medicine | 1.24 (1.03–1.50) | 0.03    | 0.89 (0.74–1.08) | 0.23    |
| TC ≥200 mg/dL or medicine | 1.29 (1.05–1.57) | 0.01    | 0.80 (0.65–0.99) | 0.04*   |
| TG ≥150 mg/dL or medicine | 1.23 (0.99–1.52) | 0.06    | 0.91 (0.74–1.12) | 0.35    |
| SBP ≥130 mmHg or medicine | 1.09 (0.87–1.36) | 0.48    | 0.98 (0.76–1.28) | 0.89    |
| DBP ≥80 mmHg or medicine | 1.06 (0.85–1.33) | 0.60    | 0.98 (0.75–1.28) | 0.86    |
| Current smoking | 1.70 (1.04–2.88) | 0.03    | 0.66 (0.36–1.22) | 0.19    |
| BMI ≥30 kg/m² | 1.00 (0.79–1.27) | 1.00 | 1.13 (0.87–1.47) | 0.37    |

| **ILI arm**     |                     |         |                    |         |
| A1C >7.0% or insulin | 1.25 (1.08–1.46) | <0.01 | 1.15 (0.95–1.39) | 0.17    |
| LDL ≥100 mg/dL or medicine | 1.15 (0.90–1.46) | 0.26 | 1.23 (0.92–1.64) | 0.17    |
| HDL ≤40 mg/dL or medicine | 1.33 (1.11–1.58) | <0.01 | 1.40 (1.12–1.75) | <0.01   |
| TC ≥200 mg/dL or medicine | 1.21 (1.00–1.48) | 0.05 | 1.28 (1.01–1.64) | 0.04    |
| TG ≥150 mg/dL or medicine | 1.75 (1.43–2.14) | <0.01 | 1.00 (0.80–1.25) | 0.98    |
| SBP ≥130 mmHg or medicine | 1.18 (0.94–1.48) | 0.16 | 1.10 (0.86–1.43) | 0.45    |
| DBP ≥80 mmHg or medicine | 1.39 (1.11–1.74) | <0.01 | 1.04 (0.78–1.37) | 0.81    |
| Current smoking | 1.57 (0.97–2.55) | 0.06 | 1.50 (0.93–2.42) | 0.10    |
| BMI ≥30 kg/m² | 1.47 (1.22–1.76) | <0.01 | 1.11 (0.90–1.38) | 0.34    |

Each row reports the statistics from a single model including terms for year, age, sex, race, education, history of CVD, duration of diabetes, prior CVD risk factor of interest status, ADM use, and BMI. Values in boldface are statistically significant. *Statistically significant in the direction opposite to hypothesis that ADM use or BMI ≥11 is associated with CVD risk factor–positive status the following year.
high sympathetic tone, and hypercholesterolemia (3,26). Of these abnormalities, we assessed only hypercholesterolemia, with mixed findings, as reported above.

We note that elevated depression screening scores are often more reflective of general emotional distress than major depressive disorder (27), and in patients with diabetes, they may reflect diabetes-related distress (28). Although depression-screening tools like the BDI have acceptable psychometric properties for detecting major depressive disorder, they often yield high rate of false-positive results (8).

All CVD risk factor measures for which we found statistically significant associations with prior ADM use, except for HDL-cholesterol, have well-documented associations with CVD morbidity and mortality. The current study does not allow us to assess the possible mechanisms that may account for the association between ADM use and subsequent CVD risk factor–positive status. Future research should assess these mechanisms:

First, individuals taking ADMs may have had a history of more severe, chronic, or recurrent depression.

Second, some individuals may have a propensity to take medicines, including ADMs and medicines that would qualify them for elevated CVD risk factor status.

Third, ADMs may contribute directly to CVD risk. TCAs are known to contribute to hyperglycemia, elevated TG levels, and weight gain (29,30). Findings on the association between TCA use and BP levels are mixed (30–32). Most SSRRs (citalopram, escitalopram, sertraline) appear to have no substantial effect on glycemia, lipid levels, BP, or weight (30). Among the SSRRs, fluoxetine has been associated with lower levels of glycemia and TG, and with weight loss, especially in the first 6 months of treatment, whereas paroxetine has been associated with the opposite effects (30). Bupropion, a noradrenaline-dopamine reuptake inhibitor, has been associated with effects on glycemia, TG levels, and weight similar to fluoxetine, and serotonin-noradrenaline reuptake inhibitors (venlafaxine, desvenlafaxine, duloxetine) and noradrenaline-serotonin specific antagonists (mirtazapine) have been associated with effects on glycemia similar to paroxetine (30). We found essentially the same pattern of associations between ADM use and subsequent CVD risk factor–positive status for all ADMs and for non-TCAs or tetracyclic ADMs. In the current study, about 80% of participants who were taking ADMs were taking non-TCAs or tetracyclic ADMs. The literature on the effects of ADMs on CVD risk factors does not provide a clear explanation for our findings. Future research should be designed to clarify the association between ADM use and CVD risk and the mechanisms that might account for this association, if any, is found.

The Markov models that we used introduced temporality into analyses to assess relationships that markers of depression in the prior year had with current CVD risk factors once the associations that prior risk factor status had with current status were controlled. Thus, the ORs that we report express the degree to which current CVD risk factor status is associated with prior markers of depression beyond what would be predicted by prior risk factor status alone. Current positive risk factor status can result from maintaining positive status from the prior year or from transitioning from prior negative status to current positive status. This approach offers insights that cannot be gained from the cross-sectional associations we previously reported (10) and strengthens the evidence for depression measures as drivers of CVD risk.

In general, our ancillary analysis findings confirmed the robustness of our original models. We found that the associations of depression symptoms and ADM use with CVD risk factors were independent of one another. We also found that the association of elevated BDI and continuous BDI with the CVD risk factors was equally strong. An analysis using log-transformed BDI scores also generated results very similar to our main analysis, as did tests for interactions of age with BDI scores and ADM use. The association between SSRI (or any non-TCAs or tetracyclic) use and CVD risk factors was essentially the same as the association between all ADM use and CVD risk factors.

The ancillary analysis that controlled for weight change during the study provided additional insight into the possible effects of elevated depression symptoms and ADM use on CVD risk factor status. This ancillary analysis reduced the number of significant associations, especially with elevated depression symptoms, but many associations, especially those with ADM use, remained significant. This suggests that factors other than weight change account for some of the associations we found and that these factors must be better understood.

**Study strengths and limitations**

This is the first study of which we are aware to simultaneously assess the independent association of two depression indicators—symptoms and ADM use—with subsequent CVD risk factor status in people with type 2 diabetes. Other study strengths include the large, multiethnic population and that depression symptoms, ADM use, and a broad range of cardiovascular risk factors were assessed systematically. Moreover, most CVD risk factors were assessed objectively rather than relying on self-reporting. Further, the design of the study, assessing depression indicators in the year prior to assessment of CVD, permitted us to draw inferences about directionality that are not possible in cross-sectional studies. The longitudinal design included assessment of depression indicators and CVD risk factors at multiple time points, enhancing the robustness of the findings.

The study also has important limitations. It was not a controlled trial assessing the effects of depression symptoms or ADM use on CVD risk factors; thus, we cannot draw definitive causal inferences. We did not study a comprehensive array of CVD risk factors, and factors such as inflammatory markers, endothelial cell dysfunction markers, and markers of kidney damage and oxidative stress were not included. Future studies of the relationship between depression indicators and CVD events will provide a more definitive picture of effects on overall CVD risk.

Another limitation is that we did not confirm that all patients took ADMs because of depression rather than for other indications, such as smoking cessation, neuropathic pain, or other psychiatric conditions, including panic disorder, social anxiety disorder, obsessive-compulsive disorder, and post-traumatic stress disorder (33). This information was not collected.

Further, we had no information about the dosage or duration of treatment with ADMs before the start of the study. Also, some participants might have failed to disclose that they were taking ADMs, whereas others who reported that they were not taking ADMs at baseline might have discontinued them very recently. However, these possibilities would mitigate the likelihood of finding an association between ADM use and ADM risk factors; therefore, they would bias the estimated associations toward the null hypothesis. Finally, we were able to identify associations only with classes of ADM, not with specific ADM.
Research and clinical implications

Future research should include more robust assessment of depression symptoms and of the reason ADMs are used (i.e., for major depressive disorder or for other conditions).

Data from the National Health and Nutrition Examination Survey, a cross-sectional survey of a nationally representative population of adults with diabetes conducted in 1999–2000, found that only 7.3% of respondents attained recommended levels of A1C, BP, and TC levels (34). If elevated depression symptoms or taking ADM add to the challenges of achieving CVD risk factor control, clinicians should be especially attentive to the fact that these patients may need more aggressive treatment to control CVD risk factors. Although the clinical relevance of the associations we report is difficult to assess, we found a dramatic increase of 24 to >50% in the odds of positive status for some CVD risk factors in study participants who had elevated depression symptoms or who were taking an ADM in the preceding year. This finding warrants serious attention.

In conclusion, among Look AHEAD participants, elevated self-reported depression symptoms and ADM use in the prior year were each independently associated with some but not all CVD risk factors during the first 4 years of the trial; significant associations for elevated risk were most consistent for ADM use. These results are consistent with the hypothesis of a potential causal link between ADM use and worsening of some CVD risk factors. Importantly, Look AHEAD will permit examination of the relationships between depression indicators and actual CVD outcomes, providing a fuller picture of the depression-CVD outcome relationship. In the meantime, more aggressive monitoring of CVD risk factors among depressed individuals and those using ADMs may be warranted.

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References

1. Egede LE, Nietert PJ, Zheng D. Depression and all-cause and coronary heart disease mortality among adults with and without diabetes. Diabetes Care 2005;28:1339–1345
2. Egede LE, Zheng D. Independent factors associated with major depressive disorder in a national sample of individuals with diabetes. Diabetes Care 2003;26:104–111
3. Lett HS, Blumenthal JA, Babiyak MA, et al. Depression as a risk factor for coronary artery disease: evidence, mechanisms, and treatment. Psychosom Med 2004;66:305–315
4. van Melle JP, de Jonge P, Spijkerman TA, et al. Prognostic association of depression following myocardial infarction with mortality and cardiovascular events: a meta-analysis. Psychosom Med 2004;66:814–822
5. Hamer M, David Batty G, Seldenrijk A, Kivimaki M. Antidepressant medication use and future risk of cardiovascular disease: the Scottish Health Survey. Eur Heart J 2011;32:437–442
6. Cohen HW, Gibson G, Alderman MH. Excess risk of myocardial infarction in patients treated with antidepressant medications: association with use of tricyclic agents. Am J Med 2000;108:2–8
7. Taylor CB, Youngblood ME, Catellier D, et al.; ENRICH-D Investigators. Effects of antidepressant medication on morbidity and mortality in depressed patients after myocardial infarction. Arch Gen Psychiatry 2005;62:792–798
8. Thombs BD, de Jonge P, Coyne JC, et al. Depression screening and patient outcomes in cardiovascular care: a systematic review. JAMA 2008;300:2161–2171
9. O’Connor CM, Jiang W, Kuchibhatla M, et al.; SADHART-CHF Investigators. Safety and efficacy of sertraline for depression in patients with heart failure: results of the SADHART-CHF (Sertraline Against Depression and Heart Disease in Chronic Heart Failure) trial. J Am Coll Cardiol 2010;56:692–699
10. Rubin RR, Knowler WC, Ma Y, et al.; Diabetes Prevention Program Research Group. Depression symptoms and antidepressant medicine use in Diabetes Prevention Program participants. Diabetes Care 2005;28:830–837
11. Rubin RR, Ma Y, Marrero DG, et al.; Diabetes Prevention Program Research Group. Elevated depression symptoms, antidepressant medicine use, and risk of developing diabetes during the diabetes prevention program. Diabetes Care 2008;31:420–426
12. Rubin RR, Gaussoin SA, Peyrot M, et al.; Look AHEAD Research Group. Cardiovascular disease risk factors, depression symptoms and antidepressant medicine use in the Look AHEAD (Action for Health in Diabetes) clinical trial of weight loss in diabetes. Diabetologia 2010;53:1581–1589
13. Ryan DH, Espeland MA, Foster GD, et al.; Look AHEAD Research Group. Look AHEAD (Action for Health in Diabetes): design and methods for a clinical trial of weight loss for the prevention of cardiovascular disease in type 2 diabetes.
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14. The Look AHEAD Research Group. Baseline characteristics of the randomized cohort for the Look AHEAD (Action for Health in Diabetes) study. Diab Vasc Dis Res 2006;3:202–215

15. Beck AT, Steers RA. Manual of the Beck Depression Inventory. San Antonio, TX, Psychological Corporation, 1993

16. American Diabetes Association. Standards of medical care in diabetes. Diabetes Care 2005;28(Suppl. 1):S4–S36

17. Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults. Executive Summary of The Third Report of The National Cholesterol Education Program (NCEP) Expert Panel on Detection, Evaluation, And Treatment of High Blood Cholesterol In Adults (Adult Treatment Panel III). JAMA 2001;285:2486–2497

18. Giorda CB, Avogaro A, Maggini M, et al.; Diabetes and Informatics Study Group. Recurrence of cardiovascular events in patients with type 2 diabetes: epidemiology and risk factors. Diabetes Care 2008;31:2154–2159

20. Diggle PJ, Liang KY, Zeger SL. Analysis of Longitudinal Data. New York, NY, Oxford University Press, 1994

21. Eeg-Olofsson K, Cederholm J, Nilsson PM, et al. New aspects of HbA1c as a risk factor for cardiovascular diseases in type 2 diabetes: an observational study from the Swedish National Diabetes Register (NDR). J Intern Med 2010;268:471–482

22. Mason PJ, Manson JE, Sesso HD, et al. Blood pressure and risk of secondary cardiovascular events in women: the Women’s Antioxidant Cardiovascular Study (WACS). Circulation 2004;109:1623–1629

23. Nordestgaard BG, Chapman MJ, Ray K, et al.; European Atherosclerosis Society Consensus Panel. Lipoprotein(a) as a cardiovascular risk factor: current status. Eur Heart J 2010;31:2844–2853

24. Guallar E, Banegas JR, Blasco-Colmenares E, et al. Excess risk attributable to traditional cardiovascular risk factors in clinical practice settings across Europe - The EURIKA Study. BMC Public Health 2011;11:704

25. McGee DL, Diverse Populations Collaboration. Body mass index and mortality: a meta-analysis based on person-level data from twenty-six observational studies. Ann Epidemiol 2005;15:87–97

26. Lederbogen F, Gilles M, Maras A, et al. Increased platelet aggregability in major depression? Psychiatry Res 2001;102:255–261

27. Coyne JC. Self-reported distress: analog or Ersatz depression? Psychol Bull 1994;116:29–45

28. Gonzalez JS, Fisher L, Polonsky WH. Depression in diabetes: have we been missing something important? Diabetes Care 2011;34:236–239

29. Aronne LJ, Segal KR. Weight gain in the treatment of mood disorders. J Clin Psychiatry 2003;64(Suppl. 8):22–29

30. McIntyre RS, Park KY, Law CWY, et al. The association between conventional antidepressants and the metabolic syndrome: a review of the evidence and clinical implications. CNS Drugs 2010;24:741–753

31. Licht CM, de Geus EJ, Seldenrijk A, et al. Depression is associated with decreased blood pressure, but antidepressant use increases the risk for hypertension. Hypertension 2009;53:631–638

32. Kemp AH, Quintana DS, Gray MA, Felmingham KL, Brown K, Gatt JM. Impact of depression and antidepressant treatment on heart rate variability: a review and meta-analysis. Biol Psychiatry 2010;67:1067–1074

33. Physicians Desk Reference. 22nd ed. Montvale, NJ, Thompson PDR, 2008

34. Saydah SH, Fradkin J, Cowie CC. Poor control of risk factors for vascular disease among adults with previously diagnosed diabetes. JAMA 2004;291:333–342