**Background:** Depressive symptoms are an established predictor of mortality and major adverse cardiac events (defined as nonfatal myocardial infarction or hospitalization for unstable angina or urgent/emergency revascularizations) in patients with acute coronary syndrome (ACS). This study was conducted to determine the acceptability and efficacy of enhanced depression treatment in patients with ACS.

**Methods:** A 3-month observation period to identify patients with ACS and persistent depressive symptoms was followed by a 6-month randomized controlled trial. From January 1, 2005, through February 29, 2008, 237 patients with ACS from 5 hospitals were enrolled, including 157 persistently depressed patients randomized to intervention (initial patient preference for problem-solving therapy and/or pharmacotherapy, then a stepped-care approach; 80 patients) or usual care (77 patients) and 80 nondepressed patients who underwent observational evaluation. The primary outcome was patient satisfaction with depression care. Secondary outcomes were depressive symptom changes (assessed with the Beck Depression Inventory), major adverse cardiac events, and death.

**Results:** At the end of the trial, the proportion of patients who were satisfied with their depression care was higher in the intervention group (54% of 80) than in the usual care group (19% of 77) (odds ratio, 5.4; 95% confidence interval [CI], 2.2-12.9 [P < .001]). The Beck Depression Inventory score decreased significantly more (t = 2.85 [P = .005]) for intervention patients (change, -5.7; 95% CI, -7.6 to -3.8; df = 155) than for usual care patients (change, -1.9; 95% CI, -3.8 to -0.1; df = 155); the depression effect size was 0.59 of the standard deviation. At the end of the trial, 3 intervention patients and 10 usual care patients had experienced major adverse cardiac events (4% and 13%, respectively; log-rank test, χ² = 3.93 [P = .047]), as well as 5 nondepressed patients (6%) (for the intervention vs nondepressed cohort, χ² = 0.48 [P = .49]).

**Conclusion:** Enhanced depression care for patients with ACS was associated with greater satisfaction, a greater reduction in depressive symptoms, and a promising improvement in prognosis.

**Trial Registration:** clinicaltrials.gov Identifier: NCT00158054

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**For editorial comment see page 585**

Patients with acute coronary syndrome (ACS) (myocardial infarction or unstable angina) who report even subsyndromal levels of depressive symptoms are at increased risk of ACS recurrence or mortality. This increased risk is observed over many years, is largely independent of other known risk factors for coronary heart disease (CHD), is strong, and has a dose-response association. The risk is particularly high for those whose depressive symptoms persist or are refractory to treatment. Although the association is not found in every study or with every ACS patient subgroup, systematic reviews, recent international data, and other accumulating research indicate that depression is a marker of increased risk of CHD events and mortality in this patient population. There have been calls for depression to be recognized as a risk marker and recommendations that patients with CHD be regularly screened for depression and be referred for treatment. However, we do not know whether patients with CHD and depressive symptoms, including many with subsyndromal symptoms, should be treated. Screening for a reliable CHD risk marker without clear evidence of how to successfully treat the risk can be problematic. In the case of depression, the suffering associated with the disorder is ar-
guably sufficient justification for treatment. Given the strength of the observational evidence, however, there have been surprisingly few trials to determine whether depression can be successfully treated in patients with ACS and the risk of ACS recurrence or mortality mitigated. The first sufficiently powered trial (Enhancing Recovery in Coronary Heart Disease [ENRICHD]; conducted in 2481 patients) to test this question found a significant but modest reduction in depressive symptoms but no mortality difference between cognitive behavioral depression therapy and usual care.17 A second trial (Myocardial Infarction and Depression–Intervention Trial [MIND-IT]; conducted in 331 patients) also found significant improvements in depression but no difference in the cardiac event rate between antidepressant treatment and usual care.18 These results were disappointing because the Sertraline Antidepressant Heart Attack Randomized Trial (conducted in 369 patients), although powered only for safety, had shown a promising trend for 6-month sertraline hydrochloride use to reduce the risk of severe cardiovascular events compared with placebo.19 Other small trials,20 and a post hoc, post-randomization responder analysis of the ENRICHD trial21 showed similar results. Given these few trials, we do not yet know whether reducing depressive symptoms improves medical prognosis in patients with ACS.

The Coronary Psychosocial Evaluation Studies (COPES) intervention trial was designed to address several reasons why previous trials may not have led to greater reductions in depressive symptoms and improvements in medical prognosis. First, the COPES trial sought to better target at-risk patients by using a 3-month observation period after ACS to eliminate patients whose symptoms spontaneously remit or respond to usual care. This strategy identifies patients with persistently elevated depressive symptoms rather than those with a diagnosis of major depressive disorder only. Second, the COPES trial adopted an approach to depression care similar to that used for the Improving Mood–Promoting Access to Collaborative Treatment (IMPACT) trial,22 including stepped care and patient preference. This approach, tailored to patients with ACS, is designed to increase the acceptance of and satisfaction with depression treatment in this population because treatment acceptance has been low in previous trials.23 We hypothesized that the COPES intervention would result in greater satisfaction with depression care and improved depressive symptoms. We also compared the rates of major adverse cardiac events (MACEs) and mortality of the depressed patients in the intervention and usual care groups with those of an observational cohort of persistently nondepressed but otherwise medically eligible patients.

**METHODS**

**RECRUITMENT, ENROLLMENT, AND INFORMED CONSENT**

Participants were recruited at 5 hospital sites (Mount Sinai Hospital and New York Presbyterian Hospital, New York, New York; and New Haven Hospital, Hospital of St Raphael, and Veterans Affairs Connecticut Healthcare System–West Haven, New Haven, Connecticut) from January 1, 2005, through February 29, 2008. Full details of the design and methods are provided elsewhere.23

Study participants were identified prospectively by monitoring hospital admissions for ACS diagnoses.24 To ensure that only patients with persistent depressive symptoms were enrolled, trial eligibility required a score of 10 or higher on the Beck Depression Inventory (BDI)25 on assessments within 1 week of hospitalization for ACS and 3 months later. Patients with BDI scores below 5 at both assessment points who met all other eligibility criteria were included in a nondepressed observational cohort.

Exclusion criteria were assessed at the hospital visit and 3-month follow-up and included alcohol or other drug dependency, dementia, current or past psychosis or bipolar disorder, terminal illness, unavailability for follow-up, BDI score of 45 or higher, or suicidality by self-report or determined during a clinical interview.

The institutional review boards at all institutions approved the protocol, and all participants provided written informed consent. To ensure equipoise, the description of the study to patients and their physicians emphasized the possible benefits and limitations of both the intervention and usual care conditions.

**RANDOMIZATION**

At each site, eligible patients were randomized on a 1:1 basis within randomly ordered blocks of 4 or 6 patients according to a table of assignments prepared in advance by the trial statistician (J.E.S.). Using a Web-based program, project coordinators specified the strata, initial, and study identification number of the person to be randomized, and the program issued the group assignment.

**INTERVENTION AND USUAL CARE PROTOCOLS**

The intervention included the following 5 essential components adapted from the IMPACT study:22 (1) an enhanced care approach, with treatment delivered by a clinical nurse specialist, psychologist, social worker, and/or psychiatrist; (2) patient choice of psychotherapy and/or pharmacotherapy; (3) a form of psychotherapy called problem-solving therapy (PST); (4) a stepped-care approach in which symptom severity was reviewed every 8 weeks and treatment was augmented according to predetermined decision rules; and (5) a standardized instrument used to track depressive symptoms.

Problem-solving therapy, as developed for the IMPACT study, has been described in detail elsewhere.22,23,26 It is a protocol driven, brief, problem focused, and designed to augment the patient’s own skills. Patients are taught how to systematically evaluate and address individual psychosocial problems. The initiation of and regular engagement in pleasant activities chosen by the patient is encouraged. Visits initially occurred weekly, in person or by telephone, with each visit lasting approximately 30 to 45 minutes. Visit frequency was decreased or increased according to the progress of individual patients and their preference.

Pharmacotherapy treatment choices included sertraline, escitalopram oxalate, venlafaxine hydrochloride, bupropion hydrochloride, and mirtazapine. A study psychiatrist or nurse practitioner prescribed appropriate medication following standard clinical practice. Intervention patients choosing pharmacotherapy were initially seen at 1- to 2-week intervals for dose titration and thereafter every 3 to 5 weeks as needed for the remainder of the 6-month trial period. If a patient was already taking an antidepressant, treatment decisions were coordinated with the prescribing physician. At the end of the trial, patients were pro-
vided with 6 further months of medication if they could not afford it but were referred to their usual care provider for follow-up. Four patients took advantage of this offer.

Steppe-care decisions for patients randomized to the intervention group were guided by responses to the 9-item Patient Health Questionnaire,27 administered at each treatment visit and formally evaluated at 8-week intervals. Patients who did not show prespecified improvement were offered the choice of switching treatments (eg, from PST to medication), adding the other treatment, or intensifying the original treatment choice, based on the treatment team’s recommendation (for details, see Burg et al23).

The control condition for the trial was usual care, as defined by the patient’s treating physicians. Physicians of the intervention and usual care patients were informed that their patients were participating in a trial and that they had elevated depressive symptoms; physicians were also told whether the patient met criteria for a major depressive episode.

DATA COLLECTION

At the time of the index ACS hospitalization, demographic, medical history, and prognostic variables were collected, including left ventricular ejection fraction and Global Registry of Acute Coronary Events (GRACE) risk score.28 At 3 months, just before randomization, a structured clinical interview (Depression Interview and Structured Hamilton questionnaire)?29 was conducted by telephone to assess the presence of a current major depressive episode and psychiatric exclusion criteria. All other measures at hospitalization, 3 months after hospitalization, and at the end of the 6-month intervention (month 9) were assessed in person. Measures at months 5 and 7 were obtained by telephone. Interviewers and those collecting medical outcome data were blinded to intervention assignment.

OUTCOME MEASURES

The primary outcome was satisfaction with depression care because previous treatments may not have been acceptable to patients with CHD.17 Patients were asked, “Over the last 2 months, how would you rate the quality of professional care you have received for your symptoms of distress or depression?” Patients responded on a 5-point Likert scale (1, excellent; 5, poor) or indicated that they had received no care for these symptoms. Depression severity was assessed by the BDI,25 a well-validated measure that is predictive of medical outcomes in this population.30,31 A BDI score of 10 or higher is consistent with at least mild to moderate depression.

For each patient-reported hospitalization, supporting documentation was gathered from the hospital record. Hospital systems were also actively surveyed for events. An end-point committee of 2 board-certified cardiologists independently reviewed and classified each hospitalization; in case of disagreement, a third board-certified cardiologist adjudicated the final end point. Cardiologists were unaware of participants’ depression or treatment status. For participants who could not be contacted or were reported deceased by a relative, the Social Security Death Index was searched to verify vital status, and death certificates were obtained. The first occurrence of a MACE (nonfatal myocardial infarction or hospitalization for unstable angina) or all-cause mortality was recorded.

ADVERSE EVENTS

Participants were asked about unanticipated problems or adverse events at each assessment (at 3, 5, 7, and 9 months) with the use of a standardized checklist covering major and minor cardiovascular symptoms and physical and psychiatric symp-

toms; these were presented regularly in a blinded fashion to the Data and Safety Monitoring Board.

STATISTICAL ANALYSIS

Differences between the intervention and usual care groups and between the trial participants and nondepressed cohort at baseline were evaluated using a $t$ test for continuous variables and $\chi^2$ analysis for categorical variables. When baseline medical covariate data were incomplete for the GRACE and Charlson indexes, a regression-based approach was used to impute the best linear predicted score based on the available items.

Outcome Analyses

Descriptive statistics based on the raw data at baseline were used to characterize the sample. Linear and nonlinear (ie, logistic) multilevel repeated-measures modeling procedures were used to generate full-information maximum-likelihood estimates of all treatment effects (outcome at 9 months or change in outcome from months 3 to 9, after the 6-month intervention). By including all subjects and all available data, this approach yields intent-to-treat estimates that are valid under the assumption that the missing data are missing at random, conditional on the observed data.20,21 Wald $\chi^2$ statistics were used to test the statistical significance of group differences at 9 months and the differential change between groups (group $\times$ time interaction). The primary outcome was the percentage of patients who rated their depression care as excellent or very good at 9 months. Change in the BDI score was a secondary outcome. Effect size was calculated as the group difference in BDI change divided by the pooled SD at baseline. Kaplan-Meier survival curves for MACES were estimated and compared using the log-rank test. All analyses were performed using SAS statistical software (version 9.2; SAS Institute Inc, Cary, North Carolina), including PROCs MIXED, NLMIXED, LIFETEST, and PHREG procedures.

Power Analysis

The 2-sided $\alpha$ was set at .05, and power was set at 0.90. The sample size was chosen to ensure this level of power to detect a 30% group difference (intervention vs usual care groups) in the proportion of patients who were satisfied with their depression care at the conclusion of the 6-month trial. This required enrolling 80 patients per group, allowing for 20% loss (eg, 64 per group with 9-month outcome data would provide a power of $=0.93$ to detect any 30% group difference in satisfaction, eg, 90% vs 60%, 65% vs 35%, or 35% vs 5%).

BASELINE CHARACTERISTICS

Patients randomized to the intervention and usual care groups were similar on all baseline variables (Table 1). In contrast, compared with those in the trial, patients in the nondepressed cohort differed on measures of depression (by definition), were less likely to be female, were more likely to be Hispanic, had more years of education, and were more likely to be married. Their index ACS was also more likely to be an ST-segment elevation or a non–ST-segment elevation myocardial infarction than unstable angina. Finally, the nondepressed cohort had significantly higher GRACE28 scores than the persistently depressed groups.
TREATMENT PREFERENCES AND INTERVENTION IMPLEMENTATION

Of the 80 patients randomized to intervention, 60 (75%) initially chose PST, 16 (20%) chose antidepressant medication, and 2 (2.5%) chose both; two patients did not state a treatment preference (Figure 1). Thirteen patients (16%) did not receive any study treatment. The mean (SD) number of treatment sessions delivered by phone ranged between 0% and 40% (mean [SD], 38.8% [36.8%]).

Of the patients who initially chose PST, 6 were additionally prescribed antidepressant medication during the course of their treatment. Of patients who initially chose antidepressant medication, 2 later additionally received PST and 1 switched to PST. Within the first 16 weeks of antidepressant treatment (during which 2 formal step reviews had been conducted by the entire depression care team), the dosage was increased once for 7 patients and twice for 2 patients. One patient’s treatment was augmented with a second antidepressant; another patient’s dosage was first decreased and later the medication was switched to another antidepressant type. Of the 2 patients who chose antidepressant treatment and PST at the beginning of the trial, 1 patient’s dosage was changed multiple times. We did not capture antidepressant increases, switches, or therapy sessions for those randomized to usual care. Overall, 7 patients terminated treatment before their study provider advised it.

PREVALENCE OF ANTIDEPRESSANT AND PSYCHOTHERAPY USE BEFORE AND AFTER THE TRIAL

Approximately one-third of the trial participants (35%) reported taking antidepressants at the 3-month randomization; at the end of the trial this was 48% in the treatment group, but remained at 30% in the usual care group (odds ratio, 4.48; 95% confidence interval [CI], 1.05-19.2 [P = .04, intention-to-treat estimates]). Participation in psychotherapy was 11% and 20%, respectively, before randomization; at the end of the trial this was 48% in the treatment group, but remained at 30% in the usual care group (odds ratio, 10.1; 95% CI, 2.32-44.3 [P = .002, intention-to-treat estimates; therefore, patient numbers not presented]).

PRIMARY TRIAL OUTCOME

The percentage of patients reporting depression care as excellent or very good at month 3 (ie, randomization) was modestly and not significantly different between groups (P = .18) (Table 2). At 9 months, however, 54% of patients in the intervention group reported this level of satisfaction with depression care compared with 19%...
in the usual care group (odds ratio, 5.4; 95% CI, 2.2-12.9 [P < .001, intention-to-treat estimates]).

SECONDARY TRIAL OUTCOMES

Depressive symptoms decreased significantly in both the intervention (mean change, −5.7; 95% CI, −7.6 to −3.8) and usual care (mean change, −1.9; 95% CI, −3.8 to −0.1) groups (Table 3). The group difference in depressive symptom decrease was also significant (mean group difference, −3.8; 95% CI, −6.5 to −1.2; t_{155} = 2.85 [P = .005]), representing a depression effect size of 0.59 (95% CI, 0.18-1.00). Table 3 also shows that the depressive symptom effects seemed to generalize across men, women, Hispanic patients, and African American patients. In an analysis of the 3-, 5-, 7- and 9-month depressive symptoms, group differences emerged 4 months into the trial (at month 7; t_{155} = 2.88 [P = .004]) and remained significant at the end of the trial (t_{155} = 2.99 [P = .003]).

Patient-reported adverse events were similar overall between the intervention and usual care groups, except that the usual care patients were significantly more likely to report experiencing a non–depression-related psychiatric problem than those in the intervention group (68 vs 59; χ^2 = 5.38 [P = .02]).

Patients in the intervention group had fewer MACE events (3 events [4%]) than did those in the usual care group (10 [13%]) or the nondepressed observational cohort (5 [6%]). Figure 2 shows the Kaplan-Meier curves for the 3 groups (log-rank test for the usual care vs intervention groups, χ^2 = 3.93 [P = .047]; for the intervention vs nondepressed group, χ^2 = 0.48 [P = .49]). No significant site differences were detected for any of the primary or secondary outcomes.
African American only (n=29)  

-1.5 (-5.3 to -2.4)  -7.9 (-12.7 to -3.1)  -6.4 (-12.6 to -0.2)  2.05 .04 

Women only (n=68)  

-2.6 (-5.1 to 0.0)  -6.5 (-9.1 to -4.0)  -4.0 (-7.6 to -0.3)  2.16 .03 

Hispanic only (n=66)  

-1.6 (-4.4 to 1.3)  -5.1 (-7.9 to -2.2)  -3.5 (-7.6 to 0.5)  1.71 .09 

Table 3. Reduction in Depressive Symptoms 3 and 9 Months After ACSa

Abbreviations: ACS, acute coronary syndrome; ellipses, not applicable.  

aData are presented as mean (95% confidence interval) unless otherwise indicated. Depressive symptom scores were assessed using the Beck Depression Inventory.25 Multilevel linear mixed models were used to estimate a group x time interaction effect of depressive symptom scores.  

In light of the damaging impact of depression on quality of life and prognosis in patients with CHD,1,30 rates of detection and effective treatment for depression remain unacceptably low in this patient population.20 It was in this spirit that the COPES trial was undertaken. The enhanced-care, patient-preference, stepped approach used herein was associated with substantial improvement in satisfaction with depression care and a significant reduction in depressive symptoms. In addition, although the study was not powered for this outcome, the intervention led to a promising difference in MACEs between randomized groups, with the MACE rate in the intervention group resembling that in the nondepressed cohort.

LIMITATIONS

First, patients selected for this trial did not include all patients with ACS. We excluded those with cognitive impairments, other life-threatening conditions, and, most important, other psychiatric conditions such as alcohol or other drug dependence and bipolar disorder. Because these comorbid conditions are highly prevalent in depressed patients, our findings might not be applicable to all patients with ACS and depressive symptoms. Second, we had a relatively small sample size, and the MACE rate was, expectedly, quite small. Thus, further trials of enhanced depression care are required to determine whether this type of treatment can improve post-ACS prognosis. Third, our patients were not blinded to their treatment status. We made every effort to blind the endpoint committee and the outcome assessors by asking patients not to reveal their group and by ensuring that assessors were not in contact with the therapist team, but this is only a single-blind trial. Fourth, we chose usual care as our control condition rather than placebo or another active control, such as clinical management. Thus, we did not account for nonspecific effects of treatment. Fifth, 13 of the 80 patients randomized to treatment never attended a first depression care visit. Another 7 terminated treatment before their care provider advised it, suggesting that, although the acceptance of our depression intervention was more than 50%, there is room for improvement. Recent studies using telephone-delivered cognitive behavioral therapy34 and combined psychotherapy with pharmacotherapy by telephone35,36 provided suggestions for novel delivery methods to further test in patients with ACS. Sixth, we did not collect cost data, which would have aided in the evaluation of this intervention. Finally, our 6-month treatment may have been too brief; we saw significant differences in depressive symptoms only after 4 months of treatment. The American College of Physicians37 recommended that clinicians continue treatment for 4 to 9 months after a satis-
factory response in patients with a first episode of major depressive disorder. For patients who have had 2 or more episodes of depression, an even longer duration of therapy may be beneficial. The depression and cardiac outcomes reported herein might be strengthened by longer depression treatment.

COMPARISON OF COPES WITH OTHER DEPRESSION INTERVENTION TRIALS IN PATIENTS AFTER ACS

Although some previous trials have shown statistically significant reductions in depressive symptoms, there were no improvements in cardiovascular outcomes.20 One possible explanation is that the depression treatment effects resulting from the modalities tested were not large enough to alter the increased risk of cardiovascular events and mortality conferred by depression.38 In fact, previous trials had 1 common finding: only clinically modest depression differences between the treatment and control groups.20 One plausible reason for this finding is that the treatments were unacceptable to patients with CHD.17 A patient’s willingness to engage in, adhere to, and continue depression treatment can determine whether the treatment succeeds or fails. Most of the depression interventions used in previous trials involving patients with CHD were originally validated with treatment-seeking outpatients with psychiatric problems; therefore, acceptance by the broader population of patients with CHD cannot be assumed.30 Previous studies have shown that fewer patients drop out of PST compared with other psychological therapies.30

Another possible explanation for the lack of improved cardiac prognosis with previously tested depression interventions is that the treatments were not sufficiently powerful.38 Recent systematic reviews of single-modality antidepressant41,42 or psychotherapy43 treatments in other patient populations showed only modest efficacy compared with placebo or usual care. Larger effect sizes have been found with multimodal44,45 or stepped-care22,46 depression treatment interventions. Until this trial, enhanced-care, stepped algorithms had not been tested in patients with CHD, but the results in other medical populations were promising.22,47-49 We thus chose to test this treatment modality in the COPES trial. We found a reasonable depression effect size (0.59) that compares favorably with those of previous interventions designed to reduce depression in patients with CHD (0.20-0.38).20

Large reductions in depressive symptoms in the control group are an issue in trials enrolling depressed patients with and without ACS.38,41 Depression is a relapsing-remitting disease30; hence, substantial reductions in symptoms and/or spontaneous remission can occur. Also, medical providers increasingly recognize depressive symptoms in patients with ACS, and some patients’ symptoms respond to the conventional depression treatment offered.21 For these reasons, we chose to include a 3-month observation period to identify patients with persistent depressive symptoms and thereby decrease the likelihood of a large reduction in depressive symptoms in the control group. We had a smaller reduction in depressive symptoms in the control group compared with other trials of depressed patients with CHD, possibly as a result of this strategy.

It is not known whether only a subset of patients who are depressed after ACS is at risk for ACS recurrence or mortality.7,51,52 We excluded more patients than we enrolled because of depressive symptom improvement, and this could be viewed as a limitation because we targeted a small sample without psychiatric diagnoses. Most observational cohort studies demonstrating depression-associated risk of ACS recurrence or mortality used a BDI score of 10 or higher to characterize depression3 rather than conventional psychiatric diagnoses.35 Participants with persistently elevated BDI scores (≥10) in these studies were found to be at risk of death.6 In the COPES trial, we similarly targeted patients with a BDI score of 10 or higher rather than just those meeting the diagnostic criteria for a psychiatric disorder. As expected with initial tests of whether reducing a risk factor offsets cardiac event rates, the impact of depression treatments on MACEs and other cardiac risks is disparate among the trials of depression treatment in patients with CHD.54 The results reported herein for the COPES trial offer promising approaches for a larger trial.

Treating depression effectively in patients with CHD may be daunting, but trials to determine the best way to manage these 2 highly prevalent and disabling diseases35 need to continue. In the secondary prevention of cardiovascular disease, stepped-care models of depression treatment with patient preference may offer an effective approach to improve depressive symptoms and satisfaction with care; whether this type of treatment can definitively improve cardiac prognosis awaits a larger trial.

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Author Contributions: Dr Schwartz had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Davidson, Rieckmann, Clemow, Schwartz, Shimbo, Kronish, Hegel, and Burg. Acquisition of data: Davidson, Clemow, Medina, Albanese, Kronish, and Burg. Analysis and interpretation of data: Davidson, Rieckmann, Clemow, Schwartz, Shimbo, Kronish, and Burg. Drafting of the manuscript: Davidson, Schwartz, Shimbo, and Burg. Critical revision of the manuscript for important intellectual content: Davidson, Rieckmann, Clemow, Schwartz, Shimbo, Medina, Albanese, Kronish, Hegel, and Burg. Statistical analysis: Schwartz and
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Correction

Error in Figure. In the article titled “Improving Prescription Drug Warnings to Promote Patient Comprehension” by Wolf et al, published in the January 11, 2010, issue of the Archives (2010;170[1]:50-56), there was an error in Figure 1. The simplified text for Label 1 should have read “Shake well before using.”