Early Cognitive Behavioral Therapy for Depression After Cardiac Surgery

Lynn V. Doering, RN, DNSc, FAAN; Belinda Chen, MPH; Rebecca Cross Bodán, PhD, NP; Marise C. Magsarili, MN, RN; Adey Nyamathi, PhD, ANP, FAAN; Michael R. Irwin, MD

Background: Despite high rates of postcardiac surgery depression, studies of depression treatment in this population have been limited. Objective: The aim of this study was to evaluate early cognitive behavioral therapy (CBT) in a home environment in patients recovering from cardiac surgery. Methods: From July 2006 through October 2009, we conducted a randomized controlled trial and enrolled 808 patients who were screened for depressive symptoms using the Beck Depression Inventory (BDI) in the hospital and 1 month later. Patients were interviewed using the Structured Clinical Interview for DSM-IV; those who met criteria for clinical depression (n = 81) were randomized to CBT (n = 45) or usual care (UC; n = 36). After completion of the UC period, 25 individuals were offered later CBT (UC + CBT). Results: Main outcomes (depressive symptoms [BDI] and clinical depression [Structured Clinical Interview for DSM-IV]) were evaluated after 8 weeks using intention-to-treat principles and linear mixed models. Compared with the UC group, in the CBT group, there was greater decline in BDI scores (β = 1.41; 95% confidence interval [CI], 0.81–2.02; P = < .001) and greater remission of clinical depression (29% [64%] vs 9% [25%]; number need to treat, 2.5; 95% CI, 1.7–4.9; P < .001). Compared with the early CBT group (median time from surgery to CBT, 45.5 days) the later UC + CBT group (median time from surgery to CBT, 122 days) also experienced a reduction in BDI scores, but the group × time effect was smaller (β = 0.79; 95% CI, 0.10–1.47; P = .03) and remission rates between the 2 groups did not differ. Conclusions: Early home CBT is effective in depressed postcardiac surgery patients. Early treatment is associated with greater symptom reduction than similar therapy given later after surgery.

KEY WORDS: cardiac surgery, cognitive behavioral therapy, depression

Depression occurs in 15% to 40% of patients who undergo cardiac surgery.1–7 After cardiac surgery, depression is a major cause of mortality; its effects are long lasting, with deaths increased for up to 10 years after surgery.8 In addition to mortality, postoperative depression has been shown to adversely affect surgical recovery and is associated with increased cardiac morbidity and decreased functional status.9

Despite the known deleterious effects of depression in this population, studies designed to treat depression early after cardiac surgery have been limited. Two studies of depression treatment after cardiac surgery have

Lynn V. Doering, RN, DNSc, FAAN
Professor and Chair, Acute Care, School of Nursing, University of California, Los Angeles.

Belinda Chen, MPH
Project Director, School of Nursing, University of California, Los Angeles.

Rebecca Cross Bodán, PhD, NP
Assistant Professor, School of Nursing, California State University, Fullerton.

Marise C. Magsarili, MN, RN
Nurse Practitioner, Kaiser Permanente Medical Center, Los Angeles, California.

Adey Nyamathi, PhD, ANP, FAAN
School of Nursing, University of California, Los Angeles.

Michael R. Irwin, MD
Professor and Director, Cousins Center for Psychoneuroimmunology, University of California, Los Angeles, Semel Institute for Neuroscience, and Department of Psychiatry and Biobehavioral Sciences, University of California, Los Angeles.

This research was supported by a grant from the National Institute of Nursing Research (NINR) (R01NR009228) to Dr Doering. The NINR had no role in the design and conduct of the study; the collection, management, analysis, and interpretation of the data; or the preparation, review, or approval of the manuscript.

DOI: 10.1097/JCN.0b013e31824d967d

Correspondence
Lynn V. Doering, RN, DNSc, FAAN, Acute Care, UCLA School of Nursing, 700 Tiverton Ave, 4-266 Factor Building, Los Angeles, CA 90095-6918 (ldoering@sonnet.ucla.edu).
been reported.\textsuperscript{10,11} Only one study has evaluated the effectiveness of cognitive behavioral therapy (CBT) in this population.\textsuperscript{10} Furthermore, novel home-based approaches to delivery of CBT for cardiac patients have not been tested systematically, with only a single pilot study of home CBT for heart failure patients reporting nonsignificant improvements in depressive symptoms when CBT was combined with an exercise component.\textsuperscript{12} Home-based therapy may be particularly advantageous for patients who are postoperative, whose recovery may limit their ability to attend traditional site-based programs. The purpose of this study was to evaluate the efficacy of CBT in a home environment among patients recovering from cardiac surgery. The primary aims of the study were to test (1) the effect of a short, 8-week course of early home CBT compared with usual care (UC) on remission of clinical depression and depressive symptom severity and (2) early home CBT compared with later home CBT on the same outcomes.

**Methods**

**Sample and Setting**

From July 2006 through October 2009, we screened 3422 patients who underwent cardiac surgery (coronary artery bypass grafting or valve replacement/repair) from 5 tertiary care hospitals in Los Angeles, California. Institutional review board approvals were obtained from all participating sites. A 2-step screening protocol was used. First, patients were excluded if they met any of the following criteria: younger than 30 years, residing outside the greater Los Angeles area, presence of cognitive impairment (Mini-Mental State Examination score < 24) or major comorbid psychiatric condition (schizophrenia, bipolar disorder, substance abuse), and autoimmune disorder or malignancy.

A second screening evaluation occurred within a month after hospital discharge. A trained research nurse called patients at home to reassess depressive symptoms. This time frame was selected because depressive symptoms at this time are more likely to be consistent with clinical depression, rather than a reactive depressive disorder or somatic symptoms related to surgery.\textsuperscript{13} Patients with Beck Depression Inventory (BDI) scores higher than 10 at either baseline or 1 month post-discharge or those with a history of depression met the second screening eligibility criteria and were interviewed at home using the Structured Clinical Interview for DSM-IV (SCID-I).\textsuperscript{14}

Inclusion criterion was a diagnosis of major or minor depression on the SCID-I. Major depression was defined as (1) presenting 5 or more symptoms (depressed mood, anhedonia, weight change, insomnia/hypersomnia, psychomotor retardation/agitation, fatigue, worthlessness/guilt, indecisiveness, thoughts of death/suicidal ideation), (2) including either depressed mood or anhedonia, (3) symptoms occurring nearly every day in a 2-week period, and (4) symptoms significant enough to interfere with daily activities. Minor depression was defined as meeting the same criteria but presenting at least 2 of the above symptoms.

**Procedures**

Of the 3422 patients, 1881 (55\%) met initial screening eligibility criteria, and 808 (43\%) consented to participate (Figure 1). Patients were approached after surgery and before hospital discharge. After consent, a brief in-hospital screening interview was conducted to assess cognitive function. Individuals with a score of 24 or higher on the Mini-Mental State Examination\textsuperscript{15} completed a brief questionnaire booklet to obtain baseline measures of sociodemographic (marital and work status) and psychobehavioral (depressive and anxiety symptoms) characteristics. Medical records were reviewed to obtain demographic data and to identify comorbidities (Charlson Comorbidity Index\textsuperscript{16}), preexisting psychiatric conditions, and current medications. Patients with preoperative depression and those on antidepressants were included in the study.

Of the 808 patients who consented to participate, 731 (90\%) were contacted for the second screening. Nineteen subjects were not interviewed because they declined the interview (n = 15), were unable to be reached (n = 2), or were excluded (n = 2). Eighty-three patients (11.4\%) met inclusion criteria for major or minor depression. Diagnoses were confirmed at consensus meetings, which included a psychiatrist (M.R.I.) and a psychologist. Two patients dropped out before randomization (1 withdrew and 1 moved out of the area), and the remaining patients (n = 81) were randomized to the UC group (n = 36) or the CBT group (n = 45) using randomization tables for each hospital created by our statistician. After randomization, 1 participant in each group was excluded when previously unidentified psychiatric conditions (ie, bipolar disorder) became evident. In addition, in the CBT group, 5 participants withdrew before starting therapy and 2 withdrew without completing therapy. Among the UC participants, after 8 weeks, 10 did not participate in later therapy because they were no longer depressed (n = 7), moved away (n = 1), or declined therapy (n = 2). Of the 23 who continued in the study, 2 withdrew without completing therapy.

The UC group received referrals to their primary care providers, along with biweekly follow-up telephone calls from study staff for 8 weeks (a total of 4 calls). Telephone calls were short (<15 minutes) and aimed primarily at maintaining ongoing contact. They followed a semiscr ipted outline, in which research staff told participants that they were “checking in with...
them” and reminded them about their enrollment in the study but did not assess depressive symptoms. Twenty-six participants of the UC group who remained depressed after the 8-week UC condition were offered CBT and comprised the later CBT (UC + CBT) group. One participant was excluded from the analysis because he requested therapy 3 years after his surgery (outlier). In the early CBT and UC groups, median time from surgery to CBT or UC was 45.5 days (interquartile range, 26.25 days). In the UC + CBT group, median time from surgery to CBT was 122.0 days (interquartile range, 29.0 days).

**Intervention**

Cognitive behavioral therapy was conducted by 4 advanced practice nurses, all of whom had expertise in

---

**FIGURE 1.** Flowchart of study participants. CBT, early cognitive behavioral therapy; SCID-I, Structured Clinical Interview for DSM-IV; UC, usual care.

---

Copyright © 2013 Wolters Kluwer Health | Lippincott Williams & Wilkins. Unauthorized reproduction of this article is prohibited.
cardiac and/or psychiatric care and completed standardized CBT training (Beck Institute, Philadelphia, Pennsylvania). Patients participated in an 8-week course of therapy, which was selected based on our pilot work, in which all patients had achieved symptom remission after 8 weeks. The average length of sessions was 50 to 60 minutes; all sessions were face to face and occurred at the patient’s home or a location of his/her choosing. All sessions were conducted 1:1 without family members present. Every effort was made to schedule weekly sessions consecutively, unless the participant requested a change. To ensure fidelity and consistency of the intervention, therapy nurses used a weekly manual, which included agenda templates and commonly used analysis forms. Agenda setting was conducted collaboratively by the nurse and patient at the beginning of each session, with the nurse typically asking “What do you want to talk about today?”. Every session included a mood check using the BDI, assignment by mutual agreement of some “homework” (ie, behavioral or cognitive tasks) to reinforce skills learned during the session. Typical topics for each session were as follows—week 1: orientation to therapy, goals, and expectation of therapy; week 2: identifying negative automatic thoughts in difficult situations during the previous week and practice reframing negative thoughts; weeks 3 to 6: continued analysis of negative thoughts, identification of core beliefs, problem solving, and coping strategies; week 7: self-therapy guide and relapse prevention; and week 8: termination, review self-therapy guide, and accomplishments of therapy. Postoperative recovery education was not the focus of therapy. However, if patients brought up concerns, such as fatigue, wound care, and diet, therapy nurses answered their questions briefly in a manner consistent with the written discharge instructions they had received and encouraged them to seek advice from their primary care providers and specialists. In addition, therapy nurses coded session activities, which reflected high levels of agreement in weekly activities, and met weekly to ensure adherence to the manual and to discuss individual cases. In case conferences, led by a study investigator (L.V.D.), fidelity to the intervention was insured through assessment and discussion of specific CBT techniques used to address common themes and problems; the study psychiatrist (M.R.I.) was available for consultation.

At the end of the initial 8 weeks, the presence of clinical depression in the CBT and UC groups was reevaluated via the SCID-I by a trained research nurse, who was blinded to the treatment group. For the UC + CBT group, SCID-I interviews were repeated at the end of therapy. A suicide protocol was in place and approved by the institutional review boards at all sites for subjects who became more depressed and expressed suicidal ideation/intention. The protocol consisted of identifying the subject’s location, devising a no-suicide plan and contract with the patient, and contacting the police and/or Psychiatric Mobile Response Team for inpatient treatment.

**Instruments**

The SCID-I was used to determine the presence or absence of clinical depression. It is widely accepted and has been used as a criterion reference in studies evaluating other interviews and questionnaires for the identification of Axis 1 disorders. The SCID-I has a reported interrater agreement of 82% and κ score of 0.79 for major depressive disorder. It was selected because it allowed screening for excluded conditions, as well as clinical depression. Criteria for major and minor depression in the SCID-I are consistent with the DSM-IV-R, as described in the inclusion criteria.

Depressive symptoms were measured with the BDI, a 21-item self-report measure used widely in cardiac patients. Used in more than 2000 empirical studies, the BDI has sound internal consistency (ie, mean Cronbach α = .82) and concurrent validity (ie, with the Hamilton Rating Scale for Depression; r = 0.75) for nonpsychiatric populations. In the current study, internal consistency of the BDI at baseline yielded an α coefficient of .87.

Anxiety, known to be highly comorbid with depression, was also assessed. It was measured by the Brief Symptom Inventory Anxiety Subscale, a 6-item questionnaire that measures psychological state anxiety symptoms. Each item is rated on a 5-point scale, with higher scores indicative of higher anxiety. The Brief Symptom Inventory has been used in related cardiac populations and has demonstrated strong internal consistency (Cronbach α = .87) and criterion validity with the Spielberger Anxiety Index (r = 0.70). Cronbach α in this study was .87.

**Analysis**

All data analyses were conducted with IBM SPSS 19 (IBM, Somers, New York, 2010). Baseline characteristics between CBT and UC groups were compared using χ² analyses and Student t test (Table 1). Comparisons between the CBT and UC groups and between the CBT and UC + CBT groups were completed using linear mixed models with maximum likelihood estimation to assess the association between BDI scores over time. A square root transformation was used on raw BDI scores to fit normality assumptions. The models included a random intercept for each subject, a fixed time factor, group, and the interaction between time and group. Analyses followed intention-to-treat (ITT) principles, in which baseline scores or last BDI scores were imputed for individuals who were excluded or did not finish 8 weeks of CBT. We tested...
First, we compared changes in the distribution of BDI analyses are reported. Second, between-group differences in the posttest presence of clinical depression was evaluated by \( \chi^2 \) analyses and ITT principles. Lastly, to determine clinical significance, we used the 2-step method of Jacobson and Truax. This method defines clinical significant change as (1) a symptom score that decreased by a reliable amount and not due to chance variation (reliable change index [RCI]) and (2) a posttreatment symptom score in the nondysfunctional range. Patients with final BDI scores lower than 10 and an RCI greater than 1.96 were considered to have experienced clinical significant change. Significance levels for all analyses were set at \( P < .05 \).

**Results**

A total of 81 patients with clinical depression were randomized (Figure 1). Forty-five individuals were assigned to CBT, and 36 were assigned to UC. There were no statistical differences between the CBT and UC groups in age, race/ethnicity, educational level, type of cardiac surgery, or marital or employment status (Table 1). However, the CBT group had fewer women than the UC group (\( P = .02 \)). There were no statistical differences in history of depression, antidepressant use, anxiety level, or other medications between the groups. Sensitivity analysis of the CBT and UC groups excluding the individuals who did not receive the allocated intervention yielded similar results. There were no statistical differences between UC participants who elected to have CBT later and those who declined or between the CBT and the UC + CBT groups.

**Cognitive Behavioral Therapy Versus Usual Care**

Participants in the CBT group had a significantly greater reduction in depressive symptoms than did those in the UC group (Table 2). Linear mixed models analyses indicated a significant group-by-time interaction effect. The BDI scores decreased over time in the CBT group, whereas BDI scores increased over time in the UC group (\( \beta = 1.41; 95\% \) confidence interval [CI], 0.81–2.02). In the posttest SCID-I interview, 29 individuals (64\%) from the CBT group experienced remission, compared with 9 (25\%) from the UC group (\( P < .001 \)) (Table 3). The number needed to produce 1 additional treatment response was 2.5 (95\% CI, 1.7–4.9).

For depressive symptom categories, the CBT group showed a significant decrease in severity of depressive symptoms, whereas the UC group did not (Figure 2).

**TABLE 1** Demographic and Clinical Characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Cognitive Behavioral Therapy Group (n = 45)</th>
<th>Usual Care Group (n = 36)</th>
<th>( P^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>63.4 (8.4)</td>
<td>63.9 (11.4)</td>
<td>.83</td>
</tr>
<tr>
<td>Body mass index, kg/m(^2)</td>
<td>30.1 (7.3)</td>
<td>31.2 (10.9)</td>
<td>.59</td>
</tr>
<tr>
<td>Charlson Comorbidity index total score</td>
<td>3.5 (2.0)</td>
<td>4.4 (2.3)</td>
<td>.07</td>
</tr>
<tr>
<td>Hospital length of stay, d</td>
<td>9.1 (5.2)</td>
<td>10.3 (6.3)</td>
<td>.33</td>
</tr>
<tr>
<td>Mini-Mental State</td>
<td>27.5 (0.9)</td>
<td>27.4 (0.9)</td>
<td>.46</td>
</tr>
<tr>
<td>Examination total score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of grafts</td>
<td>2.4 (1.4)</td>
<td>2.1 (1.3)</td>
<td>.38</td>
</tr>
<tr>
<td>Pump time, min</td>
<td>92.7 (46.1)</td>
<td>108.8 (63.7)</td>
<td>.21</td>
</tr>
<tr>
<td>Cross clamp time, min</td>
<td>65.5 (39.0)</td>
<td>87.7 (63.6)</td>
<td>.07</td>
</tr>
<tr>
<td>Ejection fraction</td>
<td>51.3 (13.8)</td>
<td>50.5 (14.7)</td>
<td>.81</td>
</tr>
<tr>
<td>Baseline BDI</td>
<td>16.8 (9.3)</td>
<td>15.1 (8.9)</td>
<td>.43</td>
</tr>
<tr>
<td>Baseline BSI</td>
<td>9.5 (7.8)</td>
<td>8.3 (6.8)</td>
<td>.54</td>
</tr>
</tbody>
</table>

Abbreviations: AVR, aortic valve replacement; BDI, Beck Depression Inventory; BSI, Brief Symptom Inventory; CABG, coronary artery bypass graft; MVR, mitral valve replacement.

\( a \) Using \( t \) tests or \( \chi^2 \) tests.
the CBT group, 14 (7%) participants had moderate to severe symptoms at baseline compared with 7 (15.6%) after CBT \((P = .001)\). In the UC group, there was no change in the distribution of symptom severity categories from baseline to 8 weeks \((P = .28)\). Twenty-two individuals (48%) in the CBT group improved across symptom categories, compared with only 8 (22%) in the UC group \((P < .001)\). Using both RCIs and a cutoff score of BDI lower than 10, we found that among 68 participants who exceeded the cutoff score at baseline, 24 (60%) of those in the CBT group experienced clinical significant improvement in symptoms compared with only 3 (11%) in the UC group \((P < .001)\) (Table 3).

### Early Cognitive Behavioral Therapy Versus Later Cognitive Behavioral Therapy

The early CBT group had a significantly greater reduction in depressive symptoms than did those in the later CBT group (UC + CBT), with a significant group-by-time interaction effect in the linear mixed models analysis (Table 2) \((\beta = 0.79; 95\% \ CI, 0.10–1.47)\). However, the magnitude of the effect was smaller than in the CBT versus UC model. Furthermore, in contrast to the CBT versus UC comparison, rates of remission of clinical depression were similar between the early CBT and UC + CBT groups \(29 \left[64.4\%\right] \text{ vs } 14 \left[56\%\right]; \ P = .33\).

For depressive symptom categories, the UC + CBT group showed no change in overall distribution of

### TABLE 2 Depressive Symptoms Across Time

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Early CBT Vs UC</th>
<th>Early CBT Vs Later CBT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>3.95 (0.97)</td>
<td>4.21 (0.32)</td>
</tr>
<tr>
<td>Time (coded 0,1)</td>
<td>-1.19 (0.20)</td>
<td>-1.85 (0.20)</td>
</tr>
<tr>
<td>Symptom duration</td>
<td>-1.59 to -0.79</td>
<td>-2.26 to -1.45</td>
</tr>
<tr>
<td>Group: early (UC = 1; CBT = 0); later (UC + CBT = 1; CBT = 0)</td>
<td>0.23 (0.28)</td>
<td>0.27 (0.47)</td>
</tr>
<tr>
<td>Group × time</td>
<td>0.81 to 2.02</td>
<td>1.01 to 1.47</td>
</tr>
<tr>
<td>-2 log likelihood</td>
<td>518.41</td>
<td>339.29</td>
</tr>
</tbody>
</table>

Abbreviations: BDI, Beck Depression Inventory; CBT, early cognitive behavioral therapy; CI, confidence interval; UC, usual care; UC + CBT, completed CBT after UC.

\(aP < .001\).

\(bP = .07\).
symptom severity categories over the 8 weeks of therapy ($P = .10$; Figure 2). Using the RCI and a cutoff score of BDI lower than 10, we found that among 20 participants who exceeded the cutoff score at baseline, only 5 (25%) of those in the UC + CBT group showed clinical significant improvement in symptoms in contrast to 24 (60%) of those in the early CBT group. Hence, more individuals in the early CBT group experienced clinical and significant changes in BDI scores compared with those in the UC + CBT group ($P = .01$; Table 3).

**Discussion**

Our findings show for the first time that a brief course of CBT delivered early after cardiac surgery by nurses in the home is effective in improving depression outcomes. Depressive symptoms were sharply reduced and clinical depression remitted more in those receiving early CBT than in those receiving UC. The decline in depressive symptoms for early CBT was clinically robust and comparable with response rates for antidepressant medications—51% for antidepressant medications versus 32% for placebo. Using the results of our posttest SCID-I interviews, early CBT produced a 64% response rate versus 25% for UC. Of note, compared with early CBT, CBT performed later (after 8 weeks of UC) was less effective in reducing depressive symptoms but yielded an equivalent rate of clinical remission.

Several factors may have accounted for the positive outcomes of early CBT treatment. The early timing of therapy may have interrupted new cases of depression at a critical time before they could become entrenched. Other investigators have shown that the duration of untreated major depressive episodes has a negative effect on treatment outcomes. It is speculated that untreated symptoms alter the character of the depression itself, yielding it more resistant to treatment. Using data from the Canadian Community Health Survey, Patten demonstrated that an individual with depressive symptoms of 3 weeks' duration has a 40% probability of recovery over the subsequent 6 weeks, whereas an individual with symptoms for 23 weeks has substantially less probability of recovery at less than 5%. Using Patten's calculator, the CBT and UC + CBT groups, with median durations of symptoms of 6.5 and 16.8 weeks, respectively, could be expected to show recovery rates of 24% and 8%, respectively. Our data exceed these predictions substantially. Our findings provide further evidence that early CBT in cardiac patients with postoperative depression is effective and provides greater symptom relief than delayed treatment does. In particular, the success of CBT may arise from its focus on cognitive, behavioral, and social dimensions of depression that may become habituated as the duration of symptoms increases.

A second explanation for our findings is that early CBT may have prevented the exacerbation of preexisting depressive episodes. This notion is consistent with the symptom relief we found, even in the absence of complete remission. Although we did not find an effect of prior history of depression on treatment outcomes, it is possible that patients who were depressed postoperatively had preexisting depressive episodes that had not been identified and were untreated or undertreated at enrollment. Further study is needed to
explicate more fully the effect of the duration of illness on early CBT outcomes after cardiac surgery. The smaller number of women enrolled in CBT compared with UC may also have influenced our findings. In the Enhancing Recovery in Coronary Heart Disease (ENRICHD) study, women did not respond to CBT as well as men did. It is possible that the greater number of women in our UC group may have contributed disproportionately to the poorer outcomes in that group. Conversely, because the CBT group included more men than women, the lack of women in that group could have contributed to their improved response to CBT. Further study with larger numbers of women in both groups is needed to fully explain the role of gender in response to CBT.

Other factors related to the use of nurses to deliver CBT could have influenced our findings. First, the nurses’ ability to use CBT techniques to guide patients in the evaluation of their postoperative medical complaints may have improved outcomes. Although this support was less comprehensive in addressing postoperative cardiac care than that in recent collaboration care models, it is possible that the nurses’ support of patient decision making regarding postoperative recovery enhanced the CBT intervention. Although the content of individual therapy sessions was driven by the patient, the CBT intervention was highly structured and focused on thoughts, emotions, and behaviors associated with depression; its structure was reinforced by specific training, monitoring of session activities, and weekly case conferences. Although it is possible that our positive outcomes were due to the attention patients received from the home visits and not the content of the therapy, a previous study of a home-based psychosocial nursing intervention in cardiac patients does not support that notion. In that study, nurses’ visits were ineffective in reducing depressive symptoms. Although those nurses were experienced in cardiac care, they received no special training in depression treatment.

Finally, patients generally trust nurses and rate their honesty and ethical standards highly. This inherent credibility may have provided a foundation for a strong therapeutic alliance and subsequent CBT effectiveness. In 2 other studies of nurse case managers, the same benefit may have accrued. However, neither study focused on psychotherapy; nurse case managers were involved primarily in coordination of antidepressant therapy. Thus, in the current report, the use of nurses for postoperative depression management with CBT is unique.

This is the first study in which nurses have conducted home CBT in patients early after cardiac surgery. In 2 previous reports of depression treatment after bypass, investigators reported positive results with CBT conducted by clinical psychologists or social workers up to 1 year after surgery and with nurse case management, which involved coordination of antidepressants and/or referral to a psychologist or psychiatrist. A third study including patients with CHD also used nurse case managers but included coordinated depression treatment and cardiac care. In the current study, compared with previous reports, therapy was initiated earlier and the role of nurses was expanded to encompass both delivery of CBT and support for routine postoperative recovery. Our findings demonstrate that nurses who have received CBT training can intervene successfully in reducing clinical depression and depressive symptoms in cardiac patients.

The current report has several limitations. Our study included a relatively small sample of depressed postcardiac surgery patients. We included only individuals who met diagnostic criteria for clinical depression. Thus, we were not able to evaluate treatment effectiveness in individuals with depressive symptoms who did not meet these criteria. We had a relatively large attrition rate, with depressed women more likely to drop out than depressed men (P < .05). Although other investigators using CBT in randomized studies have reported similar or higher rates of dropout from 30% to 37%, further study is needed to examine gender responses to therapy and to consider practical applications of CBT. Our offer of delayed treatment after completion of the UC condition may have contributed positively to retention in the UC group. Future studies may be needed to test appropriate measures to limit attrition in this depressed population. We were able to conduct posttest evaluations only and did not include long-term follow-up to evaluate the duration of study effects. Finally, we did not include a cost effectiveness analysis, which would have allowed further evaluation of the study intervention.

In summary, the current findings support early CBT for depressed postcardiac surgery patients. Although previous studies have shown that CBT is an effective treatment for depression in this population, the current findings support the unique benefit of CBT administered early in the home environment by nurses. Further study is needed to determine the long-term effects and cost effectiveness of early home CBT, to explore the relationship of symptom duration to treatment outcome after cardiac surgery, and to evaluate the use of early home CBT delivered by nurses in a wider range of cardiac patients with depressive symptoms.

### What’s New and Important
- Early home cognitive behavioral therapy (CBT) is effective in reducing depressive symptoms and clinical depression in depressed patients recovering from cardiac surgery.
- Early treatment is associated with greater symptom reduction than similar therapy given later after surgery.
- This is the first study in which nurses conducted CBT in depressed patients recovering from cardiac surgery.
Acknowledgments

We thank our collaborators at our recruitment sites: Cedars-Sinai Medical Center; Kaiser Permanente Medical Center-Sunset, Los Angeles; West Los Angeles Veterans Administration Medical Center; Long Beach Memorial Medical Care; and Ronald Reagan UCLA Medical Center. We are also grateful to all the volunteers and nurses who participated in the study.

REFERENCES


