Cognitive Therapy vs Interpersonal Psychotherapy in Social Anxiety Disorder

A Randomized Controlled Trial

Ulrich Stangier, PhD; Elisabeth Schramm, PhD; Thomas Heidenreich, PhD; Matthias Berger, MD; David M. Clark, DPhil

Context: Cognitive therapy (CT) focuses on the modification of biased information processing and dysfunctional beliefs of social anxiety disorder (SAD). Interpersonal psychotherapy (IPT) aims to change problematic interpersonal behavior patterns that may have an important role in the maintenance of SAD. No direct comparisons of the treatments for SAD in an outpatient setting exist.

Objective: To compare the efficacy of CT, IPT, and a waiting-list control (WLC) condition.

Design: Randomized controlled trial.

Setting: Two academic outpatient treatment sites.

Patients: Of 254 potential participants screened, 117 had a primary diagnosis of SAD and were eligible for randomization; 106 participants completed the treatment or waiting phase.

Interventions: Treatment comprised 16 individual sessions of either CT or IPT and 1 booster session. Twenty weeks after randomization, posttreatment assessment was conducted and participants in the WLC received 1 of the treatments.

Main Outcome Measures: The primary outcome was treatment response on the Clinical Global Impression Improvement Scale as assessed by independent masked evaluators. The secondary outcome measures were independent assessor ratings using the Liebowitz Social Anxiety Scale, the Hamilton Rating Scale for Depression, and patient self-ratings of SAD symptoms.

Results: At the posttreatment assessment, response rates were 65.8% for CT, 42.1% for IPT, and 7.3% for WLC. Regarding response rates and Liebowitz Social Anxiety Scale scores, CT performed significantly better than did IPT, and both treatments were superior to WLC. At 1-year follow-up, the differences between CT and IPT were largely maintained, with significantly higher response rates in the CT vs the IPT group (68.4% vs 31.6%) and better outcomes on the Liebowitz Social Anxiety Scale. No significant treatment × site interactions were noted.

Conclusions: Cognitive therapy and IPT led to considerable improvements that were maintained 1 year after treatment; CT was more efficacious than was IPT in reducing social phobia symptoms.

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SOCIAL ANXIETY DISORDER (SAD) is a common mental disorder that is associated with considerable vocational and psychosocial handicap and an increased risk of comorbid disorders, such as depression, other anxiety disorders, and alcohol abuse.1,2 If untreated, SAD generally takes a long-term course.3 Biological, cognitive, and interpersonal factors have been implicated in the causes of SAD,4,5 and each had led to the development of distinctive treatments. Among psychological treatments, group cognitive behavior therapies (CBTs) (Heimberg et al6 and Davidson et al7) and individual cognitive therapy (CT) have been shown to be effective. Cognitive therapy is based on the cognitive model of Clark and Wells8 of the maintenance of SAD. Efficacy has been demonstrated against exposure therapy, group CT, selective serotonin reuptake inhibitor treatment, and waiting-list control (WLC) conditions in 4 randomized controlled trials.9-12

Whereas the cognitive approach mainly emphasizes intrapersonal mechanisms, other researchers have more strongly emphasized interpersonal relationship patterns and the fulfillment of social roles in the maintenance of SAD.13 Accordingly, interpersonal psychotherapy (IPT), which was originally developed by Klorman et al14...
and Weissman et al\textsuperscript{15} for unipolar depression and which focuses on the modification of dysfunctional patterns of interpersonal relationships, may represent a useful alternative to CT. Randomized controlled trials have established that IPT is effective in depression\textsuperscript{16} and in eating disorders.\textsuperscript{17} After encouraging results in an open trial\textsuperscript{18} of patients with SAD, Lipsitz et al\textsuperscript{19} in 2008 conducted a randomized controlled trial that confirmed the improvements observed with IPT in the open trial but found no significant differences between IPT and supportive therapy.

Few direct comparisons between CBTs and IPT have been conducted. The National Institute of Mental Health Treatment of Depression Collaborative Research Program\textsuperscript{20} found that both treatments were effective, but in post hoc analysis, some evidence indicated that IPT was more effective with the most severely depressed patients. Two trials\textsuperscript{21,22} of bulimia nervosa demonstrated the superior effectiveness of CBT over IPT at the posttreatment assessment but not at the 1-year follow-up. A Norwegian group\textsuperscript{23} compared predominantly group-based versions of IPT and CT in patients with SAD in a residential setting and found limited, not significantly different, improvements of symptoms in both approaches. However, both treatments differed substantially from the individual IPT and CT programs that have received the strongest support in randomized controlled trials. Interpretation of the trial findings is further complicated by low therapist competency ratings.

The aim of the present study was to compare in SAD the short- and long-term efficacy of individual CT and IPT with that of a WLC condition. To control for therapy site allegiance effects and for capacity to deliver the treatments with a sufficient degree of competence,\textsuperscript{23,24} the investigation was conducted at 2 research centers, 1 of which (Frankfurt, Germany) had previously specialized in CT and 1 of which (Freiburg, Germany) had previously specialized in IPT. Therapists at each site were trained to provide both treatments.

**METHODS**

**DESIGN**

At each trial site, patients were randomly assigned to the CT, IPT, or WLC group. Randomization was stratified according to site and presence or absence of comorbid depression. After patient eligibility was assessed and informed consent was obtained, patients were formally enrolled in the study. Allocation was based on a computer-generated list that was concealed from the investigators. Treatment comprised up to 16 individual sessions conducted on a mainly weekly basis. A booster session was offered 2 months after the end of treatment. The WLC group received treatment after a 20-week waiting period. The main assessment points were before treatment/wait, after treatment/wait, and 1 year after treatment completion. Two treatment sites that were each experienced in conducting trials with 1 of the 2 treatment approaches participated: Frankfurt University (CT; U.S. and T.H.) and Freiburg University (IPT; E.S. and M.B.). The study design, thus, included 3 factors: (1) treatment condition (CTB vs IPT vs WLC), (2) a repeated-measures factor (pretreatment vs posttreatment vs follow-up), and (3) treatment site (Frankfurt vs Freiburg) to control for any site allegiance effects.

**PATIENTS**

Participants were recruited via the private practices of psychiatrists and psychologists, outpatient clinics, and advertisements in local newspapers and on the Internet, with use of the different referral routes varying with the local circumstances of each site. All individuals interested in participating in the study took part in a telephone screening based on the Social Phobia Inventory.\textsuperscript{25} Patients who seemed eligible were invited for a diagnostic interview. The study was approved by the ethical committees at the University of Frankfurt and the University of Freiburg. Participants were provided with a complete study description, and written consent was obtained.

Social anxiety disorder and other psychiatric diagnoses were assessed using *Structured Clinical Interview for DSM-IV* Axis I and Axis II disorders.\textsuperscript{26-28} All the diagnostic evaluations were conducted by trained and certified clinical psychologists and were reviewed by senior study investigators (U.S., E.S., and T.H.). The 17-item Hamilton Rating Scale for Depression (HRSD)\textsuperscript{29,30} was used to assess severity of depression. On the basis of 6 videotaped interviews, the intraclass correlation coefficient for the HRSD was 0.97.

Individuals were invited to participate if they met the following inclusion criteria: diagnosis of SAD according to the *DSM-IV*, any comorbid mental disorder provided that severity did not exceed that of SAD, and age 18 to 65 years. The exclusion criteria were psychosis, current substance dependency or abuse, Axis II personality disorders from the dramatic or odd cluster, severe depression (HRSD score $\geq 23$), acute suicidality, current psychopharmacologic or other psychotherapeutic treatment, and preference for psychopharmacologic treatment.

Of 697 individuals who contacted the study centers, 254 were assessed by interview; 137 individuals were excluded owing to a failure to meet the inclusion criteria or for other reasons (Figure 1). Of 44 patients who refused to participate, 8 who met the inclusion criteria withdrew after signing the consent form but before randomization. The remaining 117 individuals met the inclusion criteria and were randomized. Thirty-eight participants were allocated to CBT, 38 to IPT, and 41 to WLC. Nineteen therapists (16 clinical psychologists and 3 psychiatrists) with advanced or completed psychotherapy/clinical training participated in the trial. The 8 therapists treating patients receiving CT and 11 therapists treating patients receiving IPT had comparable levels of clinical experience (CT: 5.3 years; IPT: 6.6 years; $t_9 = -0.73$, $P = .48$), with the treatment (CT or IPT: 4.5 years; IPT: 4.1 years; $t_8 = 0.78$, $P = .44$), and experience with the treatment of SAD (CT: 1.5 years; IPT: 1.5 years; $t_9 = 0.04$, $P = .97$). In each treatment condition, therapists received 40 hours of training workshops and adhered to treatment manuals (D.M.C., unpublished data, 1997; translated and revised by Stangier, Ehlers, and Clark\textsuperscript{31}; J. D. Lipsitz, PhD, and J. C. Markowitz, PhD, unpublished data, 1996). The workshops for CT were conducted by 3 of us (U.S., T.H., and D.M.C.) and for IPT by Dr Lipsitz and one of us (E.S.). Each therapist treated at least 2 pilot cases under supervision before participating in the trial. Additional training in the form of detailed feedback on videotapes or case descriptions was provided by one of us (D.M.C.) and Dr Lipsitz. At both trial sites, continuous supervision was established for therapists in each condition. After reaching an adequate level of adherence, therapists treated an average of 4 patients each.

**TREATMENTS**

The treatments comprised 16 individual sessions conducted over 20 weeks. Most sessions were 50 minutes, but the protocol allowed therapists to extend up to 6 sessions to a maximum of...
100 minutes to facilitate behavioral experiments (CT) or in-depth discussions and role-plays (IPT). With respect to mean session length, no significant differences between both treatments (mean [SD] number of minutes per session: IPT, 65.3 [9.8]; CT, 67.8 [14.4]; t = 0.77; P = .45). Both treatments were manualized (D.M.C., unpublished data, 1997; translated and revised by Stangier, Ehlers, and Clark31; J. D. Lipsitz and J. C. Markowitz, unpublished data, 1996). Patients on the waiting list received no treatment for 20 weeks, after which they were offered 1 of the 2 treatments. None of the patients received any other form of psychotherapy or pharmacotherapy during the treatment phase of the study. The sessions were videotaped. A randomly selected subset of CT videotapes was audited by one of us (E.S.), and additional feedback was provided by Dr Lipsitz. The integrity and boundaries of each therapy were carefully monitored. Checklists of “encouraged” and “prohibited” interventions were completed by the therapist after each session to ensure that techniques unique to the other treatment were not applied.

Cognitive Therapy

The CT program was based on the cognitive model of SAD of Clark and Wells8 and included the following components8,9: (1) establishing a personal version of the model using the patient’s own thoughts, images, focus of attention, safety behaviors, and symptoms; (2) conducting role-play–based behavioral experiments to demonstrate the adverse effects of self-focused attention and safety behaviors; (3) practicing external focus of attention in nonsocial and social situations; (4) re-structuring distorted self-imagery using videotape feedback and other methods; (5) discussing surveys providing feedback on other people’s beliefs about the significance of blushing, stuttering, sweating, etc; and (6) behavioral experiments to test negative beliefs in anxiety-provoking social situations while giving up safety behaviors and adopting an external focus of attention. Therapists were instructed not to use components of IPT, such as exploring and modifying interpersonal relationships or using role-plays to enhance communication of affect and social skills.

Interpersonal Psychotherapy

For SAD, IPT was based on a revised version of the standard manual13,14 developed by Lipsitz and Markowitz (J. D. Lipsitz and J. C. Markowitz, unpublished data, 1996) and used in trials by Lipsitz et al.18,19 During the first phase of treatment, the Interpersonal Inventory is conducted with the aim of relating social anxiety symptoms to 1 of the 4 problem areas. J. Lipsitz (written communication, 2002) replaced the problem area “social deficits” with the concept of “role insecurity/role deficits” as being more specific to SAD. Most commonly used in this trial was the area of role transition, either in terms of life changes or in terms of a therapeutic role transition. Therapeutic role transition means that the patient recognizes that SAD is not part of his or her personality but rather a temporary state or role. In the second stage of treatment, the formulated problem area is addressed by clarifying roles and their associated emotions, giving advice, using role-play if indicated, and encouraging the patient to communicate and express feelings. As in standard IPT, the interventions generally aim to enable the patient to communicate and express feelings. As in standard IPT, the interventions generally aim to enable the patient to communicate and express feelings. As in standard IPT, the interventions generally aim to enable the patient to communicate and express feelings. As in standard IPT, the interventions generally aim to enable the patient to communicate and express feelings. As in standard IPT, the interventions generally aim to enable the patient to communicate and express feelings.
The primary outcome measure was treatment response as assessed by the Clinical Global Impression Scale (CGI-I). In agreement with Heimberg et al and Davidson et al, we chose CGI-I as the primary outcome measure because it is a standard primary outcome measure in psychopharmacologic studies and provides information that is of high clinical relevance. The psychometric properties of CGI-I have been found to be good. Independent assessors masked to the treatment condition completed the 7-point rating scale at the posttreatment and 1-year follow-up assessments. Patients rated 1 or 2 (markedly or moderately improved) were classified as responders, and those rated 3 or higher were classified as nonresponders.

The secondary outcome measures were independent assessor ratings on the Liebowitz Social Anxiety Scale (LSAS) and the patient-completed Social Phobia and Anxiety Inventory (SPAI) (T. Fydrich, PhD, A. Scheurich, PhD, and E. Kasten, Dipl Psych, unpublished data, 1995). Each was completed at the pretreatment/wait, posttreatment/wait, and 1-year follow-up assessments. At the end of the first session, patients rated the credibility of their treatment using a rating scale developed by Borkovec and Nau. In addition, a therapist version of this questionnaire was used to assess allegiance. After each therapy session, patients and therapists separately completed the Bernese Post-Session Report, which includes satisfactorily reliable patient- and therapist-rated therapeutic alliance scales. For the present analysis, alliance ratings after the first therapy session were used.

### STATISTICAL ANALYSES

Data were analyzed using a commercially available software package (SPSS; SPSS Inc, Chicago, Illinois). All the statistical analyses were intent-to-treat. Patients who were allocated to CT or IPT were considered to have had an adequate dose of therapy if they attended at least 12 (of 16) sessions. Individuals who attended fewer sessions were still assessed and included in the intent-to-treat analysis. Missing data were replaced using the last-observation-carried-forward approach. Categorical analyses were conducted using binary logistic regression. Dimensional measures were submitted to analyses of covariance in which pretreatment scores were controlled for. Analyses of covariance were performed separately for the posttreatment and 1-year follow-up assessments. To determine whether treatment site affected outcome, all the analyses included an estimation of site and interaction with site effect. Statistical significance was set at P < .05 (2-tailed).

### RESULTS

**DESCRIPTION OF THE SAMPLE**

Patient characteristics are given in Table 1. No significant differences were noted between treatment conditions regarding any of the sociodemographic or clinical variables. Fifty-eight percent of patients met the criteria for the generalized subtype of SAD. Fifty-four percent of patients also met the diagnostic criteria for 1 or more other current Axis I disorders: major depressive disorder (24.6%), dysthymia (13.6%), specific phobia (5.9%), and panic disorder (3.4%). Sixty-seven percent of patients met the criteria for 1 or more personality disorders, primarily avoidant type (50.8%).

**TREATMENT AND ASSESSMENT COMPLIANCE**

Figure 1 shows the flow of patients through the trial. Eleven of 76 patients (14.5%) attended fewer than 12 of 16 sessions and were considered to have received a suboptimal dose of treatment (7 patients receiving CT [18.4%] and 4 patients receiving IPT [10.5%]). Separate analyses for both sites revealed that no significant differences were noted between IPT and CT in either patient or therapist ratings of treatment credibility, treatment alliance, and adherence. For CT in either patient or therapist ratings of treatment credibility or in the quality of the therapeutic alliance. For
both treatments, credibility and therapeutic alliance scores were high. Mean (SD) treatment credibility patient ratings were as follows: IPT, 7.47 (1.26); CT, 7.82 (1.34); $F_{1,70} = 1.27, P = .26$. Mean (SD) therapist ratings were as follows: IPT, 8.2 (0.9); CT, 8.7 (0.7); $t_{17} = 1.57, P = .14$. Mean (SD) quality of the therapeutic alliance ratings by patients were as follows: CT, 1.58 (0.68); IPT, 1.45 (0.65); $F_{1,70} = 0.58, P = .45$.

Adherence to treatment protocols was assessed using the postsession therapeutic technique checklists. According to these checklists, therapists used interventions that were categorized as being unique to the relevant treatment only (CT: 54.5%, IPT: 65.5%) or that were shared by both treatments (CT: 45.4%, IPT: 35.5%). No interventions that were categorized as “prohibited” were reported.

TRIAL SITE EFFECTS

No significant site effects on the social phobia outcome measures were noted. However, there was a significant site effect on the HRSD at the posttreatment/wait assessment but not at the follow-up assessment. Irrespective of the intervention received, patients in Freiberg had higher HRSD scores at the posttreatment/wait assessment than did patients in Frankfurt. No significant treatment × site interaction effects were noted on any outcome measure at either the posttreatment/wait or the 1-year follow-up assessment. This means that the differences in outcomes among the 3 groups (CT, IPT, and WLC) reported later herein were not significantly affected by the site at which treatment was provided.

PRIMARY OUTCOME

Figure 2 shows the results for the primary outcome measure: the independent assessor ratings of treatment response using the CGI-I. At the posttreatment/wait assessment, 25 of the 38 patients (65.8%) who had undergone CT, 16 of the 38 (42.1%) who had undergone IPT, and 3 of the 41 (7.3%) in the WLC group were classified as responders. Both CT and IPT were superior to WLC (CT: Wald $\chi^2 = 21.4, P < .001$; IPT: Wald $\chi^2 = 10.55, P < .001$). In addition, CT proved superior to IPT (Wald $\chi^2 = 4.21, P = .04$). At 1-year follow-up, the difference between CT and IPT was maintained. Twenty-six of the 38 patients (68.4%) who had undergone CT and 12 of the 38 patients (31.6%) who had undergone IPT were classified as responders (Wald $\chi^2 = 9.82, P = .002$). During follow-up, significantly more patients who had received IPT sought additional psychological or pharmacologic treatment for SAD (CT, 12.1% of patients [1 psychological, 1 medication, and 2 combined treatments]; IPT, 38.2% of patients [7 psychological, 4 medication, and 2 combined treatments] ($\chi^2 = 6.03, n = 67, P = .01$). From posttreatment assessment to 1-year follow-up, 15.2% of patients receiving CT changed from nonresponse to response and 9.1% of patients from response to nonresponse. In the IPT group, these rates were 8.8% and 20.6%, respectively. Because additional treatments during follow-up may have produced further improvement, we also analyzed the proportion of patients who were classified as responders at 1-year follow-up and had not received additional treatment for SAD. The difference between CT and IPT remained significant ($P < .01$). Responder proportions were 48.6% for CT and 18.4% for IPT.

SECONDARY OUTCOMES

Table 2 provides the secondary outcome measures. At the posttreatment/wait assessment, the independent assessor ratings on the LSAS indicated that patients who received either CT or IPT showed greater improvement than did patients in the WLC group. Cognitive therapy also proved superior to IPT. The SPAI showed a similar pattern of results, although the difference between CT and IPT did not reach significance ($P = .07$). On the HRSD, the CT and IPT groups showed greater improvement than did the WLC group and did not differ from each other. At 1-year follow-up, CT remained superior to IPT on the LSAS but did not differ from IPT on the SPAI ($P = .10$) or the HRSD ($P > .40$).

EFFECT SIZES

For the primary outcome measure (CGI-I responder status), number needed to treat (NNT) is an appropriate way of quantifying effect sizes. The NNT refers to the number of patients who need to be treated with the “more effective” intervention to obtain 1 more responder as if the same number of patients had received the “less effective” intervention. For the contrasts at the posttreatment/wait assessment, NNTs were as follows: 5 for CT vs IPT, 2 for CT vs WLC, and 3 for IPT vs WLC. At 1-year follow-up, the NNT for CT compared with IPT was 3.

Table 3 provides the controlled effect sizes for the secondary outcome measures.

The results of the present study suggest that CT and IPT are effective treatments for SAD. Each treatment was associated with significantly greater improvement com-
Anxiety Inventory.

all other assessment points, the group effect is based on 2-way (treatment /H11003 SPAI, Social Phobia and Anxiety Inventory; WLC, waiting-list control. 

Abbreviations: CT, cognitive therapy; HRSD, Hamilton Rating Scale for Depression; IPT, interpersonal psychotherapy; LSAS, Liebowitz Social Anxiety Scale; SPAI, Social Phobia and Anxiety Inventory.

Table 2. Secondary Outcome Measures at the Pretreatment, Posttreatment, and Follow-up Assessmentsa

<table>
<thead>
<tr>
<th>Measure</th>
<th>Cognitive Therapy (n = 38)</th>
<th>Interpersonal Psychotherapy (n = 38)</th>
<th>Waiting-List Control (n = 41)</th>
<th>Statistic</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>LSAS score, mean (SD)</td>
<td>Pre-treatment 69.17 (23.36)</td>
<td>68.35 (22.60)</td>
<td>62.75 (26.76)</td>
<td>F2,108 = 13.86</td>
<td>.001</td>
</tr>
<tr>
<td></td>
<td>Posttreatment 39.49 (21.09)</td>
<td>48.16 (22.36)</td>
<td>59.90 (29.05)</td>
<td>F2,110 = 21.41</td>
<td>&lt;.001</td>
</tr>
<tr>
<td></td>
<td>1-y Follow-up 33.96 (20.57)</td>
<td>43.33 (25.18)</td>
<td>NA</td>
<td>F2,11 = 5.72</td>
<td>.02</td>
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<tr>
<td>HRSD score, mean (SD)</td>
<td>Pre-treatment 8.11 (5.43)</td>
<td>8.24 (5.93)</td>
<td>7.81 (6.06)</td>
<td>F2,111 = 0.12</td>
<td>.89</td>
</tr>
<tr>
<td></td>
<td>Posttreatment 5.43 (5.74)</td>
<td>4.50 (4.00)</td>
<td>8.03B (6.13)</td>
<td>F2,110 = 6.04</td>
<td>&lt;.001</td>
</tr>
<tr>
<td></td>
<td>1-y Follow-up 4.47 (5.39)</td>
<td>5.31 (4.93)</td>
<td>NA</td>
<td>F2,11 = 0.63</td>
<td>.43</td>
</tr>
<tr>
<td>SPAI score, mean (SD)</td>
<td>Pre-treatment 76.14 (16.98)</td>
<td>77.94 (15.44)</td>
<td>73.14 (23.52)</td>
<td>F2,111 = 0.06</td>
<td>.94</td>
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<tr>
<td></td>
<td>Posttreatment 51.20 (20.61)</td>
<td>59.75 (18.38)</td>
<td>69.19 (28.26)</td>
<td>F2,108 = 13.66</td>
<td>&lt;.001</td>
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<td>1-y Follow-up 49.74 (24.06)</td>
<td>55.80 (19.82)</td>
<td>NA</td>
<td>F2,10 = 2.68</td>
<td>.11</td>
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</tbody>
</table>

Abbreviations: HRSD, Hamilton Rating Scale for Depression; LSAS, Liebowitz Social Anxiety Scale (total score); NA, not applicable; SPAI, Social Phobia and Anxiety Inventory.

aWithin an assessment occasion, means with no subscript letters and those that share the same subscript letter do not differ. Means with nonoverlapping subscript letters differ at a level of at least P < .05. At the pretreatment assessment, the group effect is based on 2-way (treatment × site) analysis of variance. At all other assessment points, the group effect is based on 2-way (treatment × site) analysis of covariance, with pretreatment scores as the covariate.

Table 3. Controlled Effect Sizes at the Posttreatment and 1-Year Follow-up Assessmentsa

<table>
<thead>
<tr>
<th>Measure</th>
<th>Posttreatment Assessment</th>
<th>Follow-up Assessment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CT vs WCL</td>
<td>IPT vs WLC</td>
</tr>
<tr>
<td>LSAS</td>
<td>1.47</td>
<td>0.95</td>
</tr>
<tr>
<td>SPAI</td>
<td>1.21</td>
<td>0.79</td>
</tr>
<tr>
<td>HRSD</td>
<td>0.55</td>
<td>0.76</td>
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</table>

Abbreviations: CT, cognitive therapy; HRSD, Hamilton Rating Scale for Depression; IPT, interpersonal psychotherapy; LSAS, Liebowitz Social Anxiety Scale; SPAI, Social Phobia and Anxiety Inventory; WLC, waiting-list control.

aControlled effect sizes were computed by dividing the difference between covariance-adjusted means by the square root of the average of the variances for the groups.

pared with the WLC group. In addition, a significant advantage was found for CT over IPT on the primary outcome measure (CGI-I responder status). At posttreatment assessment, 65.8% of patients treated with CT showed marked improvement in social-phobic symptoms compared with 42.1% of those treated with IPT. At 1-year follow-up, the superiority of CT over IPT persisted, with the former showing significantly higher response rates. We additionally observed a significantly higher rate of additional nonprotocol treatment during follow-up in the IPT vs the CT group.

To ensure a valid and fair comparison of treatments, we controlled for several potential sources of bias. First, therapeutic allegiance was controlled by using 2 research sites, 1 of which had previously specialized in IPT and 1 in CT. Second, therapists at both centers received training from acknowledged experts in each treatment. Third, the therapists who provided the 2 treatments had similar levels of clinical experience and were not significantly different in their expectations for the improvements in their patients, although it cannot be excluded that slight, nonsignificant differences in outcome expectations between IPT and CT therapists might reflect different acknowledgment of empirical support for the efficacy of the treatments (at the beginning of the study, 2 controlled studies evaluating CT10 were opposed to 1 open trial of IPT18). Fourth, the quality of the therapeutic alliance, as rated by therapists and patients, was similar in the 2 treatments. Fifth, patients’ expectations after the first session were similarly high in the 2 treatments. Sixth, overall treatment compliance rates did not differ between the 2 treatments. At the site that had previously specialized in IPT, more patients in the IPT group than those in the CT group received an adequate dose of treatment, but if this were important, one would expect a higher response rate to IPT, which is the opposite of what was found. Seventh, no significant treatment × site interaction effects were noted for the primary or secondary outcome measures. Overall, it seems that there is no good reason to suppose that observed differences in outcome between CT and IPT were caused by variation in allegiance or other common, nonspecific therapy factors.

Treatment integrity was supported by providing treatment manuals developed by 1 of us (D.M.C., unpublished data, 1997; translated and revised by Stangier, Ehlers, and Clark15; and Lipsitz and Markowitz, unpublished data, 1996). In addition, 1 of us (D.M.C.) and Dr. Lipsitz conducted intensive workshops for both groups of therapists. The quality of each treatment was monitored by the primary investigators (U.S. and E.S.) and by experienced on-site supervisors. Finally, although inde-
We used the Inventory of Interpersonal Problems to evaluate self-rated interpersonal functioning. Contrary to expectations, we found no significant differences between the 2 treatments at the posttreatment assessment and significantly larger interpersonal improvements in the CT group at 1-year follow-up. A possible explanation for this result is that interpersonal problems are more likely to be resolved when the underlying dysfunctional cognitions and safety and avoidance behaviors are effectively modified.

In contrast to previous trials, we did not observe a “slower action” of IPT than of CBT. Rather, IPT could not compensate for the posttreatment differences at 1-year follow-up. Because SAD, similar to dysthymia, is a chronic disorder characterized by marked avoidance behavior, we assume that IPT may not provide sufficiently structured help (eg, exercises and homework) to overcome avoidance. Interpersonal psychotherapy was originally tailored for acute major depression. In this disorder, and possibly in bulimia nervosa, IPT might be more beneficial because acute interpersonal problems are more closely related to their etiology. In other disorders, such as dysthymia and SAD, however, interpersonal problems might be speculated to represent a consequence rather than a causative factor. Future research should investigate whether the efficacy of IPT can be increased by developing more structured interventions focusing on disorder-specific problems in SAD (eg, self-protective behaviors) or incorporate techniques that have been proved effective in CT (eg, videotape feedback).

The present study has several limitations. First, although CT was superior to IPT on the primary outcome measure (CGI-I response rate) and the other assessor ratings of social anxiety (LSAS), the self-reported social-anxiety measure (SPAI) showed only a statistical trend in favor of CT in the intent-to-treat analysis. This difference became significant in a post hoc analysis restricted to patients considered to have received an adequate dose of therapy (≥12 sessions). It is unclear why the self-report measure seems to have been less sensitive to differential treatment effects. However, this pattern of results has also been observed in some trials of pharmacologic treatments for SAD. Second, although we included patients with secondary comorbid conditions, such as mild or moderate secondary depression, the characteristics of this sample (exclusion of severe comorbid Axis I and II disorders, recruitment by advertisements, and university setting) are not fully representative of clinical practice. The generalizability of the trial findings to routine clinical care might, thus, be limited. Third, although therapist-completed ratings indicated no protocol violations, we cannot exclude the possibility that some interventions that should be unique to one treatment were occasionally used in the other treatment without being detected.

In conclusion, the results of this study demonstrate the efficacy of CT and IPT in the treatment of SAD. However, the data also provide evidence of a superiority of CT over IPT, suggesting that CT should be the preferred psychological treatment for SAD. The lack of differences between the 2 treatment sites with respect to the efficacy of CT contradicts a potential effect of allegiance and suggests that successful dissemination is possible.
For IPT, further developments might help to improve efficacy by more specifically addressing empirically supported interpersonal problems and avoidance in SAD.

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Correspondence: Ulrich Stangier, PhD, Department of Psychology, University of Frankfurt, Varrentrappstr 40-42, D-60054 Frankfurt, Germany (Stangier@psych.unifrankfurt.de)

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