Tayside–Fife clinical trial of cognitive–behavioural therapy for medication-resistant psychotic symptoms

Results to 3-month follow-up

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Background  Evidence for the efficacy of cognitive–behavioural therapy for schizophrenia is promising but evidence for clinical effectiveness is limited.

Aims  To test the effectiveness of cognitive–behavioural therapy delivered by clinical nurse specialists in routine practice.

Method  Of 274 referrals, 66 were allocated randomly to 9 months of treatment as usual (TAU), cognitive–behavioural therapy plus TAU (CBT) or supportive psychotherapy plus TAU (SPT) and followed up for 3 months.

Results  Treatment effects were modest but the CBT condition gave significantly greater improvement in overall symptom severity than the SPT or TAU conditions combined (F (1.53) = 4.14; P = 0.05). Both the CBT and SPT conditions combined gave significantly greater improvement in severity of delusions than did the TAU condition (F (1.53) = 4.83; P = 0.03). Clinically significant improvements were achieved by 7/21 in the CBT condition (33%), 3/19 in the SPT condition (16%) and 2/17 in the TAU condition (12%).

Conclusions  Cognitive–behavioural therapy delivered by clinical nurse specialists is a helpful adjunct to routine care for some people with chronic psychosis.

Declaration of interest  None.

About 30% of people with a diagnosis of schizophrenia continue to experience psychotic symptoms such as hallucinations and delusions despite treatment with antipsychotic medication (Kane, 1996). Current application of cognitive–behavioural therapy aims to help sufferers understand and manage their experience of psychosis in ways that reduce distress and interference with functioning. Meta-analytical reviews support the potential efficacy of cognitive–behavioural therapy for schizophrenia but suggest that evidence for its effectiveness in routine clinical practice is limited (Cormac et al, 2002; Pilling et al, 2002). In this paper we report a comparison of cognitive–behavioural therapy, delivered by clinical nurse specialists, with two control conditions: treatment as usual and an analytically based supportive psychotherapy condition delivered by staff in a community mental health team. Recent clinical trials of cognitive–behavioural therapy for psychosis have either not controlled for therapist contact time (Kuipers et al, 1997; Turkington et al, 2002) or have controlled for it but with the same therapists delivering both treatments (Tarrier et al, 1998; Sensky et al, 2000).

METHOD

Overview of recruitment and procedure

This study was conducted at two adjacent mental health services in Tayside and Fife (Scotland) covering a broad mix of urban and rural communities with a total catchment area of about 500 000 people. Patients were recruited by soliciting referrals from psychiatrists and psychiatric nurses working in community mental health teams, in-patient services and community care facilities. Selection criteria were as follows: patients with psychosis and a diagnosis of schizophrenia, schizoaffective disorder or delusional disorder, aged 16–65 years who are known to the psychiatric services as suffering from positive symptoms of persistent and distressing hallucinations or delusions, or both, and who have been stabilised on anti-psychotic medication for at least a 6-month period under the care of a consultant psychiatrist. Exclusion criteria were: primary diagnosis of alcoholism or drug misuse, evidence of organic brain disease and history of violence. All aspects of recruitment, screening and outcome assessment were organised and administered by an experienced psychiatrist (M.G.) over a 4-year period between January 1997 and March 2001. She had some assistance in these tasks during the latter phase of the study from an experienced community psychiatric nurse, who was trained in assessment procedures and was closely supervised.

All referrals were offered an appointment for a screening interview to establish diagnosis (using both ICD–10 (World Health Organization, 1992) and DSM–IV (American Psychiatric Association, 1994) criteria), to determine if selection criteria were met and to obtain consent. Suitable patients who consented were given further structured assessments as detailed below and included in the baseline phase of the study. They were then offered a further assessment interview 3 months later to reassess selection criteria and willingness to participate. The second screening interview was included in order to ensure stability of symptoms and as an added check on suitability for inclusion in the trial. Participants who met the criteria on both occasions were then randomised to one of three treatment conditions (cognitive–behavioural therapy plus treatment as usual – CBT; supportive psychotherapy plus treatment as usual – SPT; treatment as usual – TAU) and entered a 9-month treatment phase. They were then reassessed at the end of treatment and at a 3-month follow-up.

The randomisation procedure (sealed envelope technique) was devised by the project statistician (Cathy Hau) and administered centrally by the non-clinical project coordinator (Jen Petrie). It was carried out separately within each treatment centre using randomised permuted blocking (Johnson, 1992). Power calculations were based on the expectation of a reasonably large effect size as found at post-treatment in a clinical trial comparing a coping skills enhancement condition and a problem-solving control (Tarrier et al,
1993). This was the closest example of relevant research when the project was planned. Cohen (1992) states that having 21 patients in each group gives an 80% power of detecting a large effect size with a two-tailed significance level of 0.05. In retrospect this power calculation was not well founded because large treatment effects have not been reported generally in subsequent clinical trials. On the assumption that about two-thirds of screened patients would consent and be suitable and that about two-thirds of this number would complete, it was calculated that about 150 patients would need to be screened. In fact, these assumptions proved to be unrealistic and only about one-fifth of referrals ended up completing treatment.

Assessments

A broad range of measures were administered as part of structured interviews at initial screening, second screening, post-treatment and 3-month follow-up. Organisation and administration of the work of the independent assessors and therapists were kept strictly separate in order to maintain the blindness of the assessor. Patients also were asked not to mention any details of their treatment during post-treatment assessments, but three patients did. The primary outcome measures were chosen following perusal of the literature on assessment of psychosis (e.g. Barnes & Nelson, 1994) and consultation with clinical research teams experienced in the field. They consisted of standardised assessments of the severity of psychotic symptomatology in general, and delusions and hallucinations in particular. The former was assessed with the Positive and Negative Syndrome Scale (PANSS; Kay et al, 1987), which has been used frequently in clinical trials with psychosis and is recommended in reviews (e.g. Drake et al, 1998). There are three sub-scales (positive symptoms, negative symptoms and general psychopathology) and a total score. The Psychotic Symptom Rating Scale (PSYRATS; Haddock et al, 1999) was used to assess specific dimensions of hallucinations and delusions. The hallucination sub-scale consists of items such as frequency, duration, loudness, negative content, intensity of distress and degree of disruption. The delusion sub-scale consists of items such as amount of pre-occupation, degree of conviction, intensity of distress and disruption. Both of these instruments have good internal reliability and validity and provide a comprehensive and clinically useful picture of current mental state that should be sensitive to treatment effects. At the end of the structured interview the assessor completed the Global Assessment Scale (GAS; Endicott et al, 1976) and also Clinical Global Improvement (Guy, 1976) in terms of seven categories (marked, moderate or mild deterioration, no change, mild, moderate or marked improvement). Other self-report measures were administered to assess symptom severity, self-esteem and attitude to illness, but these are not central to the main research question and are omitted from this report.

Attitude to treatment was assessed at two time points. At the beginning and middle of therapy patients receiving either of the two psychological therapies were asked to rate the quality of the therapeutic alliance, using the Penn Helping Alliance Questionnaire (Luborsky et al., 1996), to assess their degree of improvement in terms of seven categories (much worse, moderately worse, a bit worse, unchanged, a bit better, moderately better and a lot better) and to rate the suitability of treatment and the degree to which they were learning new ways of coping. These last two measures were on 0–8 Likert scales. These data were collected on those patients receiving therapy who were given the forms to complete and managed to return them (68% in CBT, 63% in SPT). Some bias in the sample may be present. Finally, at the end of the follow-up interview, once all assessments had been completed, patients from all three treatment conditions were administered a brief semi-structured interview to assess the perceived helpfulness of treatment over the course of the trial and the quality of their relationship with their therapist/key worker. The blindness of the independent assessor was broken at this point. As a check on blindness the assessor was asked to guess treatment allocation after the final outcome assessments were completed. Analysis of these guesses found that they were no better than chance ($\chi^2=5.63, \text{d.f.}=2, \text{NS}$).

Treatment conditions

Overview

Participants in all three conditions received the usual care provided by the psychiatric services in Tayside and Fife. Services are well developed in these two areas, with a focus on community care delivered by community mental health teams. Services include regular psychiatric consultation and contact with a key worker (typically a trained community psychiatric nurse), with emergency assessment and hospital admission available as required. Facilities in the community include day care, sheltered work, supported accommodation and volunteer befriending. Specialist psychological intervention for psychosis within a cognitive–behavioural framework, although a limited resource, is offered through clinical psychology and clinical nurse specialists.

All but one of the therapists in the trial saw patients as part of their routine clinical work and it was not possible to follow a rigid protocol in respect of the duration, frequency and location of individual sessions. These aspects of the protocol were kept flexible so as to accommodate the varied needs of individual patients and therapists. The overall aim was to give each patient a maximum of 20 therapy sessions of approximately half an hour in length over a 9-month period. This level of intensity of therapy was the best that could be managed within the constraints of patient concentration and therapist workload.

Therapists

The CBT arm of the trial was delivered by five clinical nurse specialists with extensive professional experience of severe mental disorder. All had completed a recognised post-registration training in Dundee that mainly focuses on standard cognitive–behavioural therapy for common mental disorders but includes a module on psychosis. All were registered as therapists with the British Association of Behavioural and Cognitive Psychotherapy. One of these five (R.V.M.) had developed a specialist interest in cognitive–behavioural therapy for psychosis over several years and took the lead role in developing the treatment protocol, training and supervising the other therapists and treating the majority of patients. He was employed part-time on the research grant. None of the CBT therapists saw patients in the supportive psychotherapy arm of the trial, which was delivered by 16 mental health professionals (mainly nursing but also psychiatry and occupational therapy) who were attached to the clinical teams responsible for the patients referred to the trial and each saw one or two patients as part of their routine
clinical work. All had expressed an interest in developing clinical skills in psychotherapy for patients with psychosis and none had received any formal training in cognitive–behavioural therapy. They were given training and supervision by a consultant psychotherapist (L.R.T.), who has consultant responsibility for one of the day hospitals in Dundee and is director of psychotherapy training in Tayside. She took responsibility for developing the supportive psychotherapy protocol and for training and supervising the therapists. All therapists in both treatment conditions were offered bi-weekly supervision for the duration of their contact with patients in the trial, and most participated on a regular basis.

Cognitive–behavioural therapy
The treatment protocol in the CBT condition drew on best practice as exemplified by the treatment manuals of Tarrier (1992) and Kingdon & Turkington (1994). It was strongly influenced by training workshops in cognitive–behavioural therapy for psychosis delivered in Dundee by Tarrier & Turkington. The essential elements were as follows: initial emphasis on engagement, education and building a therapeutic alliance; functional analysis of key symptoms, leading to a formulation and problem list; development of a normalising rationale for the patient’s psychotic experiences; exploration and enhancement of current coping strategies; acquisition of additional coping strategies for hallucinations and delusions; and focus on accompanying affective symptomatology using relaxation training, personal effectiveness training and problem-solving as appropriate. The overall aims were: to enhance knowledge and acceptance of illness; to encourage the acquisition of specific coping skills for managing hallucinations and delusions; and to develop an understanding of personal vulnerability and how to mitigate its effects.

Supportive psychotherapy
The treatment protocol in the SPT condition was developed by L.R.T. using the framework described by Garfield in his book Unbearable Affect: A Guide to the Psychotherapy of Psychosis (Garfield, 1995). This book provides therapists with vivid case histories and concrete illustrations of therapeutic strategies that give a sense of understanding the nature of psychotic experience and the ways in which talking through these experiences can bring some measure of relief and perspective. The approach is psychodynamic in orientation (cf. De Jonghe, 1993) and seeks to understand psychotic experience as a function of being overwhelmed and unable to bear intensely charged emotional experiences. The essential elements of therapy were as follows: provision of a safe and supportive atmosphere in which to raise issues of emotional importance to the patients, with an emphasis on the non-specific factors of warmth, empathy and genuineness; opportunity for the patients to describe the narrative of their lives, including the impact of the illness, so that they can be helped to make sense of the timing of the illness and its nature and content with reference to strong and ‘unbearable’ affect regarding past aspects of personal history; and description and working through of the transference as a process through which an individual transfers onto the analyst and others, past experiences, attitudes and feelings that he or she used to experience in relation to important figures earlier in life (Bateman & Holmes, 1995).

Treatment adherence and quality
Treatment protocols in the CBT and SPT conditions required audiotaping of a random selection of early, middle and late sessions. Sessions to be taped were indicated in advance and therapists were encouraged to seek consent for recording at the beginning of treatment. In practice, tapes were not obtained from a sizeable minority of participants (38% in CBT and 35% in SPT), because either consent was refused or the therapist did not feel comfortable asking for consent from a particular patient (e.g. owing to intense paranoia), or because the quality of the recording was too poor to be usable. In total, 65 audiotapes were obtained and coded so as to conceal the therapist’s identity. Transcripts of these tapes were made and a representative sample of 45 transcripts of sessions from 23 patients were sent to an independent assessor, D.F., who used the Cognitive Therapy Scale (Vallis et al., 1986) and the Cognitive Therapy for Psychosis Adherence Scale (Startup et al., 2002) to examine treatment integrity and quality. Items that were core to both treatments were identified and additional items were included to measure key components of supportive psychotherapy. The same assessment scales were used for all transcripts and tapes.

Transcripts were necessary in order to take account of the different therapists providing treatment in the two conditions. Once the transcripts were assessed for integrity, tapes were provided to assist in the assessment of quality. It was concluded that: the two psychological therapy conditions were clearly distinct, with correct identification of therapy allocation made in 22/23 cases; adequate levels of non-specific therapy ingredients were present in both conditions, with frequently good levels of interpersonal skill; CBT sessions all involved specific, cognitive–behavioural techniques and received competent ratings on the Cognitive Therapy Scale (Vallis et al., 1986). It was also noted that only half of the SPT sessions were rated as having involved a specific psychodynamic approach. Four of thirteen CBT sessions were rated as adherent using the Cognitive Therapy for Psychosis Adherence Scale (Startup et al., 2002). This suggests that relatively few sessions included specific cognitive–behavioural therapy for psychosis techniques of the kind advocated in the Fowler et al. (1995) treatment manual. This analysis represents one of the first investigations of the nature and quality of cognitive–behavioural therapy delivered in a clinical trial with psychosis patients and points to the existence of different approaches to providing such therapy for this population.

Patient characteristics
Basic demographic characteristics of the participants are presented in Table 1. In common with other trials of this kind (e.g. Kuipers et al., 1997) the sample consisted mainly of middle-aged, single, unemployed men with a long history of illness. There were no significant differences in these characteristics across the three treatment groups. The participants as a whole were poorly educated, with 90% having left school at age 16 years or less. At the time of initial referral 85% were living in the community and 15% were in-patients in a psychiatric hospital. Of the 56 participants living in the community, 30 (54%) lived with friends or relatives, 17 (30%) lived alone and the remainder were in supported accommodation. A summary of the diagnostic and clinical status of participants is presented in Table 2.

Recruitment and attrition
A total of 274 people were referred for possible inclusion in the trial, of whom 95 (35% of initial referrals) fulfilled the initial
Fig. 1 CONSORT diagram (CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual).

Table I Demographic data on participants who entered the trial

<table>
<thead>
<tr>
<th>Variable</th>
<th>CBT (n=22)</th>
<th>SPT (n=23)</th>
<th>TAU (n=21)</th>
<th>Total sample (n=66)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male gender, n (%)</td>
<td>15 (68)</td>
<td>15 (65)</td>
<td>15 (71)</td>
<td>45 (68)</td>
</tr>
<tr>
<td>Age, years (mean ± s.d.)</td>
<td>36 (10.0)</td>
<td>37 (11.2)</td>
<td>36 (10.2)</td>
<td>36 (10.4)</td>
</tr>
<tr>
<td>Duration of illness, years (mean range)</td>
<td>15 (2–31)</td>
<td>14 (2–30)</td>
<td>10 (2–27)</td>
<td>13 (2–31)</td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Single, n (%)</td>
<td>14 (64)</td>
<td>13 (56)</td>
<td>17 (81)</td>
<td>44 (67)</td>
</tr>
<tr>
<td>Married/cohabiting, n (%)</td>
<td>5 (23)</td>
<td>5 (22)</td>
<td>4 (19)</td>
<td>14 (21)</td>
</tr>
<tr>
<td>Divorced/separated, n (%)</td>
<td>3 (14)</td>
<td>5 (22)</td>
<td>0 (0)</td>
<td>8 (12)</td>
</tr>
<tr>
<td>Employment status</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unemployed, n (%)</td>
<td>21 (96)</td>
<td>19 (83)</td>
<td>18 (86)</td>
<td>58 (88)</td>
</tr>
<tr>
<td>Sheltered work, n (%)</td>
<td>1 (4)</td>
<td>1 (4)</td>
<td>2 (9)</td>
<td>4 (6)</td>
</tr>
<tr>
<td>Employed, n (%)</td>
<td>0 (0)</td>
<td>2 (9)</td>
<td>1 (5)</td>
<td>3 (4)</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual.

criteria, entered the baseline assessment phase and were offered a further screening interview 3 months later. Of these, 66 (24% of initial referrals, 38% of 171 potentially suitable referrals) entered the study and were randomised to treatment conditions. Progress through these stages is shown in a CONSORT diagram (see Fig. 1).

Medication
Antipsychotic drug dosages over the course of the trial were available for 22 patients in CBT (100%), 19 in SPT (83%) and 18 in the TAU condition (86%). They were converted to mean daily equivalents of chlorpromazine using standard guidelines (Atkins et al., 1997) and are summarised in Table 3. The very broad confidence intervals reflect the wide variation in dosages within and between treatment conditions and none of the differences is significant. The increases in dosage post-treatment were a result of increasing use of atypical antipsychotics. Four of the fifteen patients who were started on an atypical were prescribed clozapine. No relationship was found between outcome and commencement of atypical antipsychotics. In order to assess whether or not outcome could be attributed to changes in medication, following randomisation the sample was grouped according to dose change (increased, unchanged, decreased) and an analysis of variance was conducted on difference scores on the primary outcome measure (total PANSS score) from baseline to post-treatment and baseline to follow-up. No significant differences were found at either post-treatment (F (2,55)=0.63, P=0.54) or follow-up (F (2,52)=1.51, P=0.23). The largest decreases in symptom severity were associated with reductions in medication dose.

RESULTS

Statistical analysis
Comparative efficacy of treatment conditions on the main outcome measures (PANSS, PSYRATS, GAS) was analysed first in terms of changes in mean scores over time, using an average of the baseline scores for each patient as the pre-treatment measure. A second analysis examined the proportion of patients in each treatment condition who showed at least 25% and 50% decreases in symptom severity on the PANSS and PSYRATS at post-treatment and follow-up. Both figures were chosen in order to make meaningful comparisons with the three main clinical trials published in this area, one of which (Kuipers et al., 1997) used a measure of clinically significant change that broadly equates with the lower figure, whereas the other two (Tarrier et al., 1998; Sensky et al., 2000) reported outcomes in terms of the higher figure. Although both figures are arbitrary, most clinicians with experience of chronic schizophrenia would probably regard a 25% improvement as being worthwhile and a 50% improvement as representing an important clinical change. Comparisons were made of the ratings of overall improvement across treatment conditions, from the perspective of both patient and assessor, and of the patient’s overall

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shown in Table 4. It can be seen that base-

Mean scores across treatment conditions

Response to medication

Partial response

16 (73)  
17 (74)  
17 (81)

Poor response

6 (27)  
6 (26)  
4 (19)

Significant medical history

None

11 (50)  
15 (65)  
12 (60)

Present

9 (41)  
5 (22)  
7 (35)

Alcohol/substance misuse

2 (9)  
3 (13)  
1 (5)

Table 2 Clinical status at initial screening

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>CBT (n=22)</th>
<th>SPT (n=23)</th>
<th>TAU (n=21)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F20, schizophrenia</td>
<td>18 (82)</td>
<td>23 (100)</td>
<td>18 (86)</td>
</tr>
<tr>
<td>F22, delusional disorder</td>
<td>1 (4)</td>
<td>–</td>
<td>1 (5)</td>
</tr>
<tr>
<td>F25, schizoaffective disorder</td>
<td>3 (14)</td>
<td>–</td>
<td>2 (9)</td>
</tr>
</tbody>
</table>

Response to medication

Partial response

16 (73)  
17 (74)  
17 (81)

Poor response

6 (27)  
6 (26)  
4 (19)

Significant medical history

None

11 (50)  
15 (65)  
12 (60)

Present

9 (41)  
5 (22)  
7 (35)

Alcohol/substance misuse

2 (9)  
3 (13)  
1 (5)

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual.

Table 3 Changes in prescribed antipsychotic drugs

<table>
<thead>
<tr>
<th>Chlorpromazine equivalents, mg/day (mean (95% CI))</th>
<th>CBT (n=22)</th>
<th>SPT (n=19)</th>
<th>TAU (n=18)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st screening</td>
<td>691 (441–942)</td>
<td>711 (522–901)</td>
<td>575 (302–849)</td>
</tr>
<tr>
<td>2nd screening</td>
<td>604 (392–816)</td>
<td>747 (527–967)</td>
<td>630 (333–927)</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>833 (512–1155)</td>
<td>1021 (542–1501)</td>
<td>865 (303–1427)</td>
</tr>
<tr>
<td>Follow-up</td>
<td>627 (371–882)</td>
<td>961 (486–1437)</td>
<td>911 (343–1478)</td>
</tr>
</tbody>
</table>

Patients changing total antipsychotic drug doses at follow-up evaluation, n (%)

<table>
<thead>
<tr>
<th>Reduced</th>
<th>12 (54)</th>
<th>7 (37)</th>
<th>3 (17)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased</td>
<td>6 (27)</td>
<td>7 (37)</td>
<td>7 (39)</td>
</tr>
</tbody>
</table>

Patients changing antipsychotic drug prescription at follow-up evaluation, n (%)

<table>
<thead>
<tr>
<th>Discontinued</th>
<th>1 (4)</th>
<th>2 (10)</th>
<th>0 (0)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No change</td>
<td>6 (27)</td>
<td>7 (37)</td>
<td>7 (39)</td>
</tr>
<tr>
<td>Started during study</td>
<td>6 (27)</td>
<td>6 (32)</td>
<td>3 (17)</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual.

attitude to treatment at follow-up and during therapy. All analyses reported were conducted on available data using SPSS version 10 for Windows. There was a relatively small amount of missing data at post-treatment (9%) and follow-up (14%). The analyses were repeated with the missing values replaced either with previous values carried forward or with group means, and the same pattern of significance was found.

Changes in severity from baseline

Mean scores across treatment conditions for the four main outcome measures are shown in Table 4. It can be seen that baseline scores are very stable for all measures across all three treatment conditions. An average of the baseline scores was used as the pre-treatment measure. Repeated measures analyses of variance were first conducted with three levels of treatment (CBT v. SPT v. TAU) and three time points (baseline, post-treatment, follow-up). There were significant effects for time for all variables except the GAS but no significant time × treatment interaction effects or contrasts for any of the measures. The analyses were repeated with two levels of treatment, first to provide a more powerful test of CBT effects against an aggregate of the two control conditions (CBT v. SPT and TAU) and, second, to test for the effects of receiving psychological therapy (CBT and SPT v. TAU). The first set of analyses replicated the initial analysis with the exception of the total PANSS score, where there was a significant time × treatment within-subject effect (F(2,106)=3.15, P=0.047). The linear effect of the time × treatment interaction was also significant (F(1,53)=4.14, P=0.047). The degree of overall improvement was greater in the CBT condition than in the other two conditions combined. The second set of analyses again replicated the initial analysis with the exception of the PSYRATS delusions score, where there was a non-significant effect of time (F(2,106)=2.79, P=0.06) but a significant time × treatment within-subject effect (F(2,106)=3.25, P=0.043). The linear effect of time × treatment interaction was also significant (F(1,53)=4.83, P=0.032). The degree of overall improvement in severity of delusions was significantly greater for those patients receiving psychological therapy (either CBT or SPT) than TAU.

In order to test for the location and magnitude of change in the two outcome measures with significant time × treatment interactions (PANSS total score and PSYRATS delusions), difference scores were calculated between average baseline scores and post-treatment and follow-up scores. The significance of these differences was examined with paired t-tests and the results are summarised in Table 5. It can be seen that the one significant change on the PANSS total score occurs in the CBT condition at follow-up. The changes at post-treatment are generally much smaller and none is significant. Changes on the PSYRATS delusions sub-scale are of a similar magnitude in the CBT and SPT conditions, with small non-significant changes at post-treatment and larger, significant changes at follow-up. Corresponding changes in TAU are very small and non-significant at post-treatment and follow-up.

Clinically significant improvement

Table 6 summarises the proportion of patients in each treatment condition who showed 25% and 50% improvements on PANSS total scores. The right-hand part of the table shows the difference in these proportions for patients treated with CBT relative to those with either SPT or TAU, from which the ‘number needed to treat’ statistic can be calculated. At post-treatment it can be seen that no patients showed a 50% reduction and relatively small proportions (10–20%) showed a
Table 4  Mean scores (s.d.) on main outcome measures rated by independent assessor

<table>
<thead>
<tr>
<th></th>
<th>CBT</th>
<th>SPT</th>
<th>TAU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n  Mean</td>
<td>s.d.</td>
<td>n  Mean</td>
</tr>
<tr>
<td>PANSS total score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st screening</td>
<td>22 101.2</td>
<td>14.7</td>
<td>23 96.3</td>
</tr>
<tr>
<td>2nd screening</td>
<td>22 101.2</td>
<td>14.7</td>
<td>23 95.0</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>22 96.2</td>
<td>17.7</td>
<td>19 95.2</td>
</tr>
<tr>
<td>Follow-up</td>
<td>21 87.0</td>
<td>23.1</td>
<td>19 93.5</td>
</tr>
<tr>
<td>PSYRATS delusions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st screening</td>
<td>22 13.9</td>
<td>5.3</td>
<td>22 12.5</td>
</tr>
<tr>
<td>2nd screening</td>
<td>22 14.1</td>
<td>4.5</td>
<td>23 12.3</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>22 13.3</td>
<td>5.4</td>
<td>19 11.8</td>
</tr>
<tr>
<td>Follow-up</td>
<td>21 11.1</td>
<td>5.8</td>
<td>19 9.7</td>
</tr>
<tr>
<td>PSYRATS hallucinations</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st screening</td>
<td>21 23.7</td>
<td>11.4</td>
<td>23 24.1</td>
</tr>
<tr>
<td>2nd screening</td>
<td>21 23.0</td>
<td>11.3</td>
<td>23 23.6</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>21 17.9</td>
<td>13.2</td>
<td>19 20.6</td>
</tr>
<tr>
<td>Follow-up</td>
<td>20 18.5</td>
<td>12.8</td>
<td>19 18.0</td>
</tr>
<tr>
<td>Global Assessment Scale</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st screening</td>
<td>21 32.6</td>
<td>6.2</td>
<td>23 32.5</td>
</tr>
<tr>
<td>2nd screening</td>
<td>22 32.0</td>
<td>4.8</td>
<td>22 34.9</td>
</tr>
<tr>
<td>Post-treatment</td>
<td>21 33.2</td>
<td>7.7</td>
<td>17 33.8</td>
</tr>
<tr>
<td>Follow-up</td>
<td>18 35.8</td>
<td>9.7</td>
<td>12 36.3</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual; PANSS, Positive and Negative Syndrome Scale; PSYRATS, Psychotic Symptom Rating Scale.

25% reduction. Although twice as many patients in the CBT condition achieved a 25% improvement, the absolute numbers are small and the differences are not significant. Thus, the number of patients that would need to be treated with CBT in order to achieve a difference of this kind, relative to SPT and TAU, is now six and five, respectively. Only four patients in the study achieved a 50% decrease in overall symptomatology, three of whom were in the CBT condition.

Patient and assessor ratings of overall improvement
Judgements of overall improvement over the course of the trial were made by the independent assessor at the 3-month follow-up.

For the patients as a whole, 10% were rated as having some degree of deterioration, 40% as being unchanged and 50% as having improved to some degree. When broken down by treatment condition there was a trend for a greater proportion of patients in the CBT condition being rated as improved to some degree (63%) compared with SPT (36%) and TAU (50%), and also for a larger proportion of CBT patients (15%) to be rated as having deteriorated to some degree in comparison with SPT (6%) and TAU (6%) patients. Judgements of overall improvement made by patients at the 3-month follow-up were broadly similar, although a rather greater proportion rated themselves as being worse (18%), a rather smaller proportion as unchanged (19%) and a rather larger proportion as being better to some degree (64%). None of the differences between treatment conditions was significant.

Table 5  Mean change (95% CI) from baseline at post-treatment and 3-month follow-up on outcome measures with significant overall treatment x time effects

<table>
<thead>
<tr>
<th></th>
<th>CBT</th>
<th>SPT</th>
<th>TAU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean (95% CI)</td>
<td>t</td>
<td>d.f.</td>
</tr>
<tr>
<td>PANSS total score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-treatment</td>
<td>5.1 (–0.3 to 10.5)</td>
<td>2.0</td>
<td>21</td>
</tr>
<tr>
<td>3-month follow-up</td>
<td>14.0 (3.5 to 24.5)</td>
<td>2.8</td>
<td>20</td>
</tr>
<tr>
<td>PSYRATS delusions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Post-treatment</td>
<td>0.7 (–0.9 to 2.4)</td>
<td>0.9</td>
<td>21</td>
</tr>
<tr>
<td>3-month follow-up</td>
<td>2.9 (0.7 to 5.1)</td>
<td>2.7</td>
<td>20</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual; PANSS, Positive and Negative Syndrome Scale; PSYRATS, Psychotic Symptom Rating Scale.
context that virtually identical ratings were made of the quality of the therapeutic relationship when measured by a sub-scale of the Penn Helping Alliance during the early stage of therapy. This rating was completed by 15 patients in CBT and 12 patients in SPT on a 25-point scale with a minimum of 5 and a maximum of 30 (mean score CBT = 25.1, s.d. = 3.0, mean score SPT = 25.7, s.d. = 3.0).

Difference in perceived helpfulness of therapy may have been due to differences in ratings of perceived suitability of treatment made by patients in CBT (n = 13) in comparison with patients in SPT (n = 12). At the first administration of this rating (0–8 scale: 0, not at all suitable; 8, very suitable) there were no significant differences between treatments (mean score CBT = 5.5, s.d. = 2.1; mean score SPT = 5.0, s.d. = 2.0), but at the second administration, during the middle stage of therapy, mean ratings of suitability of treatment by patients in CBT were significantly higher than those by patients in SPT (CBT = 6.6, s.d. = 1.6; SPT = 4.9, s.d. = 2.1; t = 2.27, d.f. = 23, P = 0.033). There was a similar (although non-significant) trend with regard to ratings of learning new ways of coping with problems and difficulties.

**DISCUSSION**

**Summary of findings**

The study provides evidence that cognitive–behavioural therapy in particular and psychological therapy in general give some additional benefit in overall symptom reduction when added to routine care. About one-third of patients in the CBT arm showed a 25% reduction in overall symptomatology. No evidence was found for treatment effects for auditory hallucinations but the CBT and SPT conditions combined produced significantly greater reductions in the severity of delusions than did TAU. Symptomatic improvement following psychological treatment was apparent at follow-up rather than at post-treatment, in line with the findings of a recent meta-analytical review (Pilling et al, 2002), and was not associated with changes in overall functioning as assessed by the GAS. From a clinical perspective, the overall reductions in symptomatology were relatively modest but it should be noted that the severity of initial symptoms in the trial and the magnitude of change in the CBT condition, as measured by the PANSS, are comparable with the figures reported in a recent clinical trial of the efficacy of atypical neuroleptics in chronic schizophrenia (Volavka et al, 2002). Notwithstanding symptom change, satisfaction with treatment was generally positive across all three treatment conditions, and expressions of ‘definite satisfaction’ with treatment were significantly higher in the CBT condition (70%) than in SPT (37%) or TAU (2%).

**Methodological issues**

This clinical trial is one of the first investigations of the efficacy of cognitive–behavioural therapy for psychosis in which treatment was delivered as part of routine clinical practice. The following features have helped to ensure methodological rigour: a 3-month pre-treatment baseline, to ensure the stability of the presenting symptomatology; outcome evaluation with standardised measures by an independent assessor, an experienced psychiatrist, blind to treatment allocation at post-treatment and 3-month follow-up; a credible supportive psychotherapy condition carried out by separate therapists to control for the non-specific effects of CBT; a TAU condition to control for changes over time; an independent evaluation of adherence to psychological treatment protocols and of the quality of therapy delivered; and a research team drawn from nursing, psychiatry and clinical psychology with varying professional commitments to

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**Table 6** Proportion of patients showing greater than 25% and 50% improvement on PANSS total scores at post-treatment and follow-up with absolute benefit increase (ABI) and number needed to treat (NNT)

<table>
<thead>
<tr>
<th>Patients showing improvement, n (%)</th>
<th>Rate of symptom reduction because of CBT</th>
<th>Relative to SPT</th>
<th>Relative to TAU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CBT</td>
<td>SPT</td>
<td>TAU</td>
</tr>
<tr>
<td>Post-treatment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PANSS total score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥ 25%</td>
<td>4 (18)</td>
<td>2 (11)</td>
<td>2 (11)</td>
</tr>
<tr>
<td>≥ 50%</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Follow-up</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PANSS total score</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≥ 25%</td>
<td>7 (33)</td>
<td>3 (16)</td>
<td>2 (12)</td>
</tr>
<tr>
<td>≥ 50%</td>
<td>3 (14)</td>
<td>1 (5)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual; PANSS, Positive and Negative Syndrome Scale.

**Table 7** Patients’ attitudes to treatment received over the course of the trial, assessed at 3-month follow-up

<table>
<thead>
<tr>
<th></th>
<th>CBT</th>
<th>SPT</th>
<th>TAU</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Was the treatment you received a positive, helpful experience?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No, definitely not</td>
<td>1 (5%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>No, not really</td>
<td>–</td>
<td>3 (18%)</td>
<td>1 (5%)</td>
</tr>
<tr>
<td>Yes and no</td>
<td>3 (15%)</td>
<td>3 (18%)</td>
<td>1 (5%)</td>
</tr>
<tr>
<td>Yes, to some extent</td>
<td>2 (10%)</td>
<td>4 (25%)</td>
<td>10 (60%)</td>
</tr>
<tr>
<td>Yes, definitely</td>
<td>14 (70%)</td>
<td>6 (37%)</td>
<td>5 (30%)</td>
</tr>
<tr>
<td>How well did you get on with your therapist or main contact?</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Not at all well</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Not very well</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>So-so</td>
<td>–</td>
<td>2 (12%)</td>
<td>–</td>
</tr>
<tr>
<td>Reasonably well</td>
<td>10 (50%)</td>
<td>7 (41%)</td>
<td>8 (50%)</td>
</tr>
<tr>
<td>Extremely well</td>
<td>10 (50%)</td>
<td>8 (47%)</td>
<td>8 (50%)</td>
</tr>
</tbody>
</table>

CBT, cognitive–behavioural therapy plus treatment as usual; SPT, supportive psychotherapy plus treatment as usual; TAU, treatment as usual.
cognitive–behavioural therapy, psycho-
dynamic therapy and biological psychiatry.

Methodological limitations include:
lower than expected numbers in each
treatment condition as a consequence of a
higher than expected proportion of patients
refusing to participate at the initial screen-
ing (44% of 171 suitable referrals); variable
medication regimes between and within
treatment conditions following baseline
assessment, producing potential biases with
the small sample sizes; and a potential
confound of treatment condition with
differences in therapist experience and
expertise in delivering psychological
therapy for patients with chronic psychosis.
The advantage of CBT over SPT may be
a function of the considerable experience
and expertise in delivering psychological
therapy with this population developed
by clinical nurse specialists in cognitive–
behavioural therapy over several years. In
particular, the construction and present-
ation of a formulation – a potentially
powerful intervention with this popu-
lation – takes considerable practice. Ther-
apists in the SPT arm were, on the whole,
less experienced and found the role of
therapist more challenging.

Comparison with other trials
Three other randomised controlled trials
have been published that specifically test
the efficacy of cognitive–behavioural ther-
apy for chronic psychosis: the London–East
Anglia study (Kuipers et al, 1997, 1998),
the Manchester Wellcome study (Tarrier
et al, 1998) and the London–Newcastle
Wellcome study (Sensky et al, 2000). In
common with the present investigation, all
three trials found evidence that cognitive–
behavioural therapy was more effective
than alternatives and associated with low
drop-out rates and high patient satisfaction.
However, the proportion of patients in
cognitive–behavioural therapy judged to
be treatment responders at follow-up was
found to be 65% in the London–East
Anglia study (based on 25% reductions in
symptom severity), 33% in the Manchester
Wellcome study (based on 50% reductions
in symptom severity) and 63% in the
London–Newcastle Wellcome study (also
based on 50% reductions in symptom severity).
Varying operational definitions of a
clinically significant treatment response,
as well as different measures, timescales
and selection procedures, make direct
comparisons across studies problematic
but it would appear that all three of these
trials found higher rates of treatment
responders than the 33% found in the
present study (based on 25% reductions
in symptom severity).

Variable nature of cognitive–
behavioural therapy for psychosis
One possible explanation for this apparent
difference in outcome may lie in the type
of treatment delivered. The analysis of
treatment integrity and quality completed
by D.F. found that the style of cognitive–
behavioural therapy delivered was com-
petent with respect to the standard treatment
approach used but included few of the spe-
cific adaptations for psychosis advocated in
The variations in practice within cogni-
tive–behavioural therapy for psychosis raise
the possibility that some approaches may
be more effective than others, and this
is an important area for future research.
Cognitive–behavioural therapy is a de-
manding therapy with complex presenta-
tions (Durham et al, 2000) and there is
much scope for better models and illustra-
tions of good practice. Clinicians working
with cognitive–behavioural therapy and
psychosis do not have the benefit of a range
of videotape material demonstrating
therapeutic styles and techniques but
instead have to work from treatment
manuals containing general guidelines and
brief transcripts. It is not an easy task to
apply this knowledge to the wide variety
of clinical presentations of psychosis and
it is often hard to know how an ‘expert’ in
the field might handle the difficulties that
arise. On the other hand, the present study
demonstrates that significant benefits in
routine clinical practice can result from
using standard approaches to cognitive–
behavioural therapy for psychotic disorder
and that the therapy can be of value even
with therapists given relatively brief
training (Turkington et al, 2002).

Whatever the explanation for the
apparent discrepancy in outcomes between
the present study and previous clinical
trials, it will be important to conduct
further studies in the field to clarify the
relationship between the nature and quality
of therapy delivered and clinically signifi-
cant outcomes. The current debate on this
issue tends to be polarised between those
who believe that cognitive–behavioural
therapy is clearly efficacious and those
who seem sceptical of any incorporation
of specialised psychological intervention
into routine care. Our own view of the evi-
dence is that people with chronic psychosis
should be given the opportunity to engage
in systematic psychological therapy on the
grounds that most will find this of some
value and a few will be able to use the
opportunity to make a significantly better
adjustment for the future.

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Behaviour Therapy for People with Psychosis: A Clinical


