

Cochrane Database of Systematic Reviews

Supportive therapy for schizophrenia (Review)

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Buckley LA, Maayan N, Soares-Weiser K, Adams CE. Supportive therapy for schizophrenia. *Cochrane Database of Systematic Reviews* 2015, Issue 4. Art. No.: CD004716. DOI: 10.1002/14651858.CD004716.pub4.

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[Intervention Review]

Supportive therapy for schizophrenia

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Editorial group: Cochrane Schizophrenia Group.

Publication status and date: Edited (no change to conclusions), published in Issue 11, 2017.

Citation: Buckley LA, Maayan N, Soares-Weiser K, Adams CE. Supportive therapy for schizophrenia. *Cochrane Database of Systematic Reviews* 2015, Issue 4. Art. No.: CD004716. DOI: 10.1002/14651858.CD004716.pub4.

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ABSTRACT

Background

Supportive therapy is often used in everyday clinical care and in evaluative studies of other treatments.

Objectives

To review the effects of supportive therapy compared with standard care, or other treatments in addition to standard care for people with schizophrenia.

Search methods

For this update, we searched the Cochrane Schizophrenia Group's register of trials (November 2012).

Selection criteria

All randomised trials involving people with schizophrenia and comparing supportive therapy with any other treatment or standard care.

Data collection and analysis

We reliably selected studies, quality rated these and extracted data. For dichotomous data, we estimated the risk ratio (RR) using a fixed-effect model with 95% confidence intervals (CIs). Where possible, we undertook intention-to-treat analyses. For continuous data, we estimated the mean difference (MD) fixed-effect with 95% CIs. We estimated heterogeneity (I² technique) and publication bias. We used GRADE to rate quality of evidence.

Main results

Four new trials were added after the 2012 search. The review now includes 24 relevant studies, with 2126 participants. Overall, the evidence was *very low quality*.

We found no significant differences in the primary outcomes of relapse, hospitalisation and general functioning between supportive therapy and standard care.

There were, however, significant differences favouring other psychological or psychosocial treatments over supportive therapy. These included hospitalisation rates (4 RCTs, n = 306, RR 1.82 CI 1.11 to 2.99, *very low quality of evidence*), clinical improvement in mental state (3 RCTs, n = 194, RR 1.27 CI 1.04 to 1.54, *very low quality of evidence*) and satisfaction of treatment for the recipient of care (1 RCT, n = 45, RR 3.19 CI 1.01 to 10.7, *very low quality of evidence*). For this comparison, we found no evidence of significant differences for rate of relapse, leaving the study early and quality of life.



When we compared supportive therapy to cognitive behavioural therapy CBT), we again found no significant differences in primary outcomes. There were very limited data to compare supportive therapy with family therapy and psychoeducation, and no studies provided data regarding clinically important change in general functioning, one of our primary outcomes of interest.

Authors' conclusions

There are insufficient data to identify a difference in outcome between supportive therapy and standard care. There are several outcomes, including hospitalisation and general mental state, indicating advantages for other psychological therapies over supportive therapy but these findings are based on a few small studies where we graded the evidence as *very low quality*. Future research would benefit from larger trials that use supportive therapy as the main treatment arm rather than the comparator.

PLAIN LANGUAGE SUMMARY

Supportive therapy for schizophrenia

Schizophrenia is a severe mental illness with 'positive symptoms' such as hallucinations (hearing voices and seeing things) and delusions (having strange beliefs). People with schizophrenia also suffer from disorganisation and 'negative symptoms' (such as tiredness, apathy and loss of emotion). People with schizophrenia may find it hard to socialise and find employment. Schizophrenia is considered one of the most burdensome illnesses in the world. For some people it can be a lifelong condition. Most people with schizophrenia will be given antipsychotic medications to help relieve the symptoms. In addition to this they can also receive therapy, of which there are various types.

One therapy often given to people with schizophrenia is supportive therapy, where typically after a person is established in the care of mental health services, they will receive general support rather than specific talking therapies such as cognitive behavioural therapy (CBT). For example, in consultations with health professionals there will often be time given to listening to people's concerns, providing encouragement, or even arranging basic help with daily living. Many people with schizophrenia also receive support from their family and friends. Supportive therapy has been described as the treatment of choice for most people with mental illness and may be one of the most commonly practiced therapies in mental health services.

It is, however, difficult to answer the question of exactly what supportive therapy is. It is difficult to find a widely accepted definition of supportive therapy. For the purposes of this review, supportive therapy includes any intervention from a single person aimed at maintaining a person's existing situation or assisting in people's coping abilities. This includes interventions that require a trained therapist, such as supportive psychotherapy, as well as other interventions that require no training, such as 'befriending'. Supportive therapy does not include interventions that seek to educate, train or change a person's way of coping.

The aim of this review is to assess the effectiveness of supportive therapy compared to other specific therapies or treatment as usual. This update is based on a search run in 2012; the review now includes 24 randomised studies with a total of 2126 people. The studies compared supportive therapy either with standard care alone or a range of other therapies such as CBT, family therapy and psychoeducation. The participants continued to receive their antipsychotic medication and any other treatment they would normally receive during the trials. Overall, the quality of evidence from these studies was very low. There is not enough information or data to identify any real therapeutic difference between supportive therapy and standard care. There are several outcomes, including hospitalisation, satisfaction with treatment and general mental state, indicating advantages for other psychological therapies over supportive therapy. However, these findings are limited because they are based on only a few small studies where the quality of evidence is very low. There was very limited information to compare supportive therapy with family therapy and psychoeducation as most studies in this review focused on other psychological therapies, such as CBT. Apart from one study presenting data on death, there was no information on the adverse effects of supportive therapy. In summary, there does not seem to be much difference between supportive therapy, standard care and other therapies. Future research would benefit from larger studies where supportive therapy is the main treatment.

Ben Gray, Senior Peer Researcher, McPin Foundation: http://mcpin.org/



Summary of findings for the main comparison. SUPPORTIVE THERAPY versus STANDARD CARE for schizophrenia

SUPPORTIVE THERAPY versus STANDARD CARE for schizophrenia

Patient or population: patients with schizophrenia

Settings: inpatients and outpatients Intervention: SUPPORTIVE THERAPY

Comparison: STANDARD CARE

Outcomes			Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	- (33 /0 Ci)	(studies)	(GRADE)	
	Control	SUPPORTIVE THERAPY versus STANDARD CARE				
Global state: Relapse Follow-up: 2 years	321 per 1000	309 per 1000 (141 to 678)	RR 0.96 (0.44 to 2.11)	54 (1 study)	⊕⊕⊝⊝ very low ^{1,2,3}	
Service outcomes: Hospitalisation Follow-up: 6 months	42 per 1000	42 per 1000 (3 to 628)	RR 1 (0.07 to 15.08)	48 (1 study)	⊕⊙⊙ very low ^{1,2,3}	
Mental state: no clinically important improvement Follow-up: 1 to 2 years	898 per 1000	853 per 1000 (736 to 997)	RR 0.95 (0.82 to 1.11)	98 (2 studies)	⊕⊝⊝⊝ very low ^{1,2,3}	
Leaving the study early Follow-up: 10 weeks to 2 years	166 per 1000	143 per 1000 (88 to 232)	RR 0.86 (0.53 to 1.4)	354 (4 studies)	⊕⊝⊝⊝ very low ^{1,2,4}	
General functioning GAS Follow-up: 1 years		The mean general functioning in the intervention groups was 1.4 higher (5.09 lower to 7.89 higher)		29 (1 study)	⊕⊝⊝⊝ very low ^{1,2,5,6}	
Satisfaction with treatment: Recipient of care not satisfied with treatment Follow-up: 1 years	238 per 1000	436 per 1000 (179 to 1000)	RR 1.83 (0.75 to 4.47)	44 (1 study)	⊕⊝⊝⊝ very low ^{1,2,5}	

Quality of life

WBS

The mean quality of life in the inter-260 ⊕⊝⊝⊝ vention groups was (1 study) very low 1,2,5,6 **2.73 lower**

*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** Risk ratio;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

(6.04 lower to 0.58 higher)

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

- ¹ Imprecision: serious relatively few participants were included and few events; confidence intervals are wide.
- ² Publication bias: strongly suspected four studies or fewer reported data for this outcome.
- ³ Risk of bias: serious one study had an unclear risk of bias for randomisation, allocation concealment and blinding.
- ⁴ Risk of bias: serious one study had an unclear risk of bias for randomisation and allocation concealment; two studies had an unclear risk for blinding.
- ⁵ Risk of bias: serious one study had an unclear risk of bias for blinding.
- ⁶ Indirectness: serious we wanted to collect binary data for this outcome, however, only a proxy scale measure was available.

Summary of findings 2. SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT for schizophrenia

SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT for schizophrenia

Patient or population: patients with schizophrenia

Settings: inpatients and outpatients **Intervention: SUPPORTIVE THERAPY**

Comparison: ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants (studies)	Quality of the evidence	Comments
	Assumed risk				(GRADE)	
		SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREAT- MENT				
Global state: Relapse Follow-up: 2 to 3 years	435 per 1000	517 per 1000 (287 to 939)	RR 1.19 (0.66 to 2.16)	270 (5 studies)	$\oplus \circ \circ \circ$ very low 1,2,3,4	

Service outcomes: Hospitalisation Follow-up: 12 weeks to 2 years	122 per 1000	222 per 1000 (135 to 365)	RR 1.82 (1.11 to 2.99)	306 (4 studies)	⊕⊙⊝⊝ very low ^{3,4,5}	
Mental state: No clinically important improvement Follow-up: 1 to 2 years	594 per 1000	754 per 1000 (618 to 915)	RR 1.27 (1.04 to 1.54)	194 (3 studies)	⊕⊝⊝⊝ very low ^{2,3,4,6}	
Leaving the study early Follow-up: 10 weeks to 3 years	249 per 1000	257 per 1000 (217 to 302)	RR 1.03 (0.87 to 1.21)	1412 (19 studies)	⊕⊕⊕⊝ moderate ⁷	
General functioning GAF and GAS Follow-up: 12 to 18 months	See comment	See comment	Not estimable	78 (2 studies)	⊕⊙⊙ very low 3,8,9,10	There was very high hetero- geneity for this outcome so data were not pooled. ¹¹
Satisfaction with treatment: Recipient of care not satisfied with treatment Follow-up: 1 years	136 per 1000	435 per 1000 (138 to 1000)	RR 3.19 (1.01 to 10.07)	45 (1 study)	⊕⊝⊝ very low ^{4,12,13}	
Quality of life QLS		The mean quality of life in the intervention groups was 0.07 lower (21.11 lower to 20.97 higher)		19 (1 study)	⊕⊙⊙ very low 4,10,13,14	

*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** Risk ratio;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹ Risk of bias: serious - four studies had an unclear risk of bias for randomisation and allocation concealment. All studies had an unclear risk of bias for blinding of participants and two had a high risk of bias for blinding of outcome assessments.

² Inconsistency: serious - there was high heterogeneity for this outcome.

³ Imprecision: serious - relatively few participants were included and few events; confidence intervals are wide.

⁴ Publication bias: strongly suspected - four studies or fewer reported data for this outcome.

⁵ Risk of bias: serious - three studies had an unclear risk of bias for randomisation and allocation concealment. All studies had an unclear risk of bias for blinding of participants and two were unclear for blinding of outcome assessments.

6 Risk of bias: serious - one study had an unclear risk of bias for randomisation. All studies had an unclear risk of bias for blinding of participants and one had an unclear risk of bias for blinding of outcome assessments.

- 7 Risk of bias: serious 13 studies had an unclear risk of bias for randomisation and 14 for allocation concealment. Fifteen studies had an unclear risk of bias for blinding of participants and one had a high risk of bias. Three studies two had a high risk of bias for blinding of outcome assessments and eight were unclear. One study had a high risk of bias for incomplete outcome data, and in three it was unclear.
- 8 Risk of bias: serious one study had an unclear risk of bias for allocation concealment. Both studies had an unclear risk of bias for blinding of participants and outcome assessments.
- ⁹ Inconsistency: very serious there was high heterogeneity for this outcome and data were not pooled.
- ¹⁰ Indirectness: serious we wanted to collect binary data for this outcome, however, only a proxy scale measure was available.
- 11 One study found no difference in general functioning on the GAS, the other study found a difference in favour of supportive therapy on the GAF.
- ¹² Risk of bias: serious the study had an unclear risk of bias for blinding of participants and outcome assessors.
- ¹³ Imprecision: very serious relatively few participants were included and few events; confidence intervals are very wide.
- ¹⁴ Risk of bias: serious the study had an unclear risk of bias for blinding of participants and outcome assessments.

Summary of findings 3. SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY for schizophrenia

SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY for schizophrenia

Patient or population: patients with schizophrenia

Settings: inpatients and outpatients **Intervention: SUPPORTIVE THERAPY**

Comparison: COGNITIVE BEHAVIOURAL THERAPY

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	- (33 /0 Ci)	(studies)	(GRADE)	
	Control	SUPPORTIVE THERAPY ver- sus COGNITIVE BEHAVIOURAL THERAPY				
Global state: Relapse Follow-up: 2 to 3 years	429 per 1000	471 per 1000 (219 to 1000)	RR 1.1 (0.51 to 2.41)	181 (4 studies)	⊕⊝⊝⊝ very low ^{1,2,3,4}	
Service outcomes: Hospitalisation Follow-up: 12 weeks to 18 months	156 per 1000	249 per 1000 (132 to 468)	RR 1.6 (0.85 to 3)	153 (3 studies)	⊕⊝⊝⊝ very low ^{3,4,5}	
Mental state: No clinically important improvement Follow-up: 1 to 2 years	594 per 1000	731 per 1000 (529 to 1000)	RR 1.23 (0.89 to 1.70)	194 (3 studies)	⊕⊝⊝⊝ very low ^{2,4,6}	
Leaving the study early Follow-up: 10 weeks to 3 years	150 per 1000	158 per 1000 (116 to 217)	RR 1.05 (0.77 to 1.44)	812 (12 studies)	⊕⊕⊕⊝	

					moderate ⁷	
General functioning GAF and GAS	See comment	See comment	Not estimable	78 (2 studies)	⊕⊙⊙ very low 3,4,8,9,10	There was very high heterogeneity for this outcome so data were not pooled. ¹¹
Satisfaction with treatment: Recipient of care not satisfied with treatment Follow-up: 1 years	136 per 1000	435 per 1000 (138 to 1000)	RR 3.19 (1.01 to 10.07)	45 (1 study)	⊕⊙⊙ very low ^{4,12,13}	
Quality of life RSES Follow-up: 12 weeks		The mean quality of life in the intervention groups was 1.7 lower (5.19 lower to 1.79 higher)		65 (1 study)	⊕⊙⊙ very low ^{3,4,10}	

^{*}The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

- 1 Risk of bias: serious three trials had an unclear risk of bias for randomisation and allocation concealment. All studies had an unclear risk for blinding of participants, while two studies were rated as high risk for blinding of outcome assessors.
- ² Inconsistency: serious there was high heterogeneity for this outcome.
- ³ Imprecision: serious relatively few participants were included and few events; confidence intervals are wide.
- ⁴ Publication bias: strongly suspected four studies or fewer reported data for this outcome.
- ⁵ Risk of bias: serious one study had an unclear risk of bias for randomisation and two for allocation concealment. All were unclear risk for blinding of participants and two for blinding of outcome assessors.
- 6 Risk of bias: serious one study had an unclear risk of bias for randomisation. All three had an unclear risk of blinding of participants and one for blinding of outcome assessors.
- 7 Risk of bias: serious seven studies had an unclear risk of bias for randomisation and allocation concealment. Nine studies had an unclear risk of bias for blinding of participants. Two studies two had a high risk of bias for blinding of outcome assessments and four were unclear.
- 8 Risk of bias: serious one study had an unclear risk of bias for allocation concealment. Both studies had an unclear risk of bias for blinding of participants and outcome assessments.
- ⁹ Inconsistency: very serious there was high heterogeneity for this outcome and data were not pooled.
- ¹⁰ Indirectness: serious we wanted to collect binary data for this outcome, however, only a proxy scale measure was available.
- 11 One study found no difference in general functioning on the GAS, the other study found a difference in favour of supportive therapy on the GAF.



¹³ Imprecision: very serious - relatively few participants were included and few events; confidence intervals are very wide.

Summary of findings 4. SUPPORTIVE THERAPY versus FAMILY THERAPY for schizophrenia

SUPPORTIVE THERAPY versus FAMILY THERAPY for schizophrenia

Patient or population: patients with schizophrenia

Settings: inpatients and outpatients Intervention: SUPPORTIVE THERAPY

Comparison: FAMILY THERAPY

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk	(== :: = :,	(studies)	(GRADE)	
	Control	SUPPORTIVE THERAPY versus FAMILY THERA- PY				
Global state: Relapse Follow-up: 2 to 3 years	See comment	See comment	Not estimable	87 (2 studies)	⊕⊝⊝⊝ very low ^{1,2,3,4}	There was high heterogeneity for this outcome and data were not pooled. ⁵
Service outcomes: Hospitalisation Follow-up: 2 years	300 per 1000	579 per 1000 (267 to 1000)	RR 1.93 (0.89 to 4.17)	39 (1 study)	⊕⊝⊝⊝ very low ^{3,4,6}	
Mental state: Episode of affective symptoms	292 per 1000	499 per 1000 (239 to 1000)	RR 1.71 (0.82 to 3.6)	48 (1 study)	⊕⊝⊝⊝ very low ^{3,4,7,8}	
Leaving the study early	182 per 1000	231 per 1000 (102 to 525)	RR 1.27 (0.56 to 2.89)	87 (2 studies)	⊕⊙⊙⊙ very low ^{1,3,4}	
General functioning: No paid work	600 per 1000	576 per 1000 (342 to 978)	RR 0.96 (0.57 to 1.63)	39 (1 study)	⊕⊝⊝⊝ very low ^{3,4,6,9}	
Satisfaction with treatment - not reported	See comment	See comment	Not estimable	-	See comment	No studies reported data for satisfaction with treatment.

Quality of life - not reported

See comment See comment

Not estimable

See comment

*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** Risk ratio;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

- 1 Risk of bias: very serious both studies had an unclear risk of bias for randomisation, allocation concealment and blinding of participants. One study was rated high risk of bias for blinding of outcome assessors.
- ² Inconsistency: very serious there was high heterogeneity for this outcome and data were not pooled.
- ³ Imprecision: serious relatively few participants were included and few events; confidence intervals are wide.
- ⁴ Publication bias: strongly suspected two studies or fewer reported data for this outcome.
- ⁵ One study found no difference in relapse rates, the other study found a difference in favour of family therapy.
- ⁶ Risk of bias: serious the study had an unclear risk of bias for randomisation, allocation concealment and blinding of participants.
- 7 Risk of bias: very serious the study had an unclear risk of bias for randomisation, allocation concealment and blinding of participants. It was rated high risk of bias for blinding of outcome assessors.
- ⁸ Indirectness: serious we wanted to collect binary data for no clinical improvement in general symptoms, however, only a proxy measure was available.
- ⁹ Indirectness: serious we wanted to collect binary data for no overall improvement in general functioning, however, only a proxy measure was available.

Summary of findings 5. SUPPORTIVE THERAPY versus PSYCHOEDUCATION for schizophrenia

SUPPORTIVE THERAPY versus PSYCHOEDUCATION for schizophrenia

Patient or population: patients with schizophrenia

Settings: inpatients and outpatients Intervention: SUPPORTIVE THERAPY

Comparison: PSYCHOEDUCATION

Outcomes	Illustrative comparative risks* (95% CI)	Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk Corresponding risk	(33 /0 Ci)	(studies)	(GRADE)	
	Control SUPPORTIVE THERAPY versu PSYCHOEDUCATION				

Global state: Relapse - not reported	See comment	See comment	Not estimable	-	See comment	No studies re- ported data on relapse.
Service outcomes: Hospitalisation Follow-up: 6 months	87 per 1000	42 per 1000 (4 to 429)	RR 0.48 (0.05 to 4.93)	47 (1 study)	⊕⊝⊝⊝ very low ^{1,2,3}	
Mental state. No clinically important improvement Follow-up: 6 months	600 per 1000	966 per 1000 (576 to 1000)	RR 1.61 (0.96 to 2.68)	19 (1 study)	⊕⊝⊝⊝ very low ^{2,3,4}	
Leaving the study early Follow-up: 6 months	250 per 1000	142 per 1000 (52 to 385)	RR 0.57 (0.21 to 1.54)	71 (2 studies)	⊕⊝⊝⊝ very low ^{2,3,5}	
General functioning - not reported	See comment	See comment	Not estimable	-	See comment	No studies re- ported data for general func- tioning.
Satisfaction with treatment - not reported	See comment	See comment	Not estimable	-	See comment	No studies reported data for satisfaction with treatment.
Quality of life QLS		The mean quality of life in the intervention groups was 0.07 lower (21.11 lower to 20.97 higher)		19 (1 study)	⊕⊝⊝⊝ very low ^{2,3,4,6}	

^{*}The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RR: Risk ratio;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

¹ Risk of bias: serious - the study had an unclear risk of bias for randomisation, allocation concealment and blinding of participants and outcome assessors.

² Imprecision: very serious - relatively few participants were included and few events; confidence intervals are wide.

³ Publication bias: strongly suspected - two studies or fewer reported data for this outcome.

⁴ Risk of bias: serious - the study had an unclear risk of bias for randomisation, allocation concealment and blinding of participants.



⁵ Risk of bias: serious - both studies had an unclear risk of bias for randomisation, allocation concealment and blinding of participants and one study was unclear risk for outcome assessors.

⁶ Indirectness: serious - we wanted to collect binary data for this outcome, however, only a proxy scale measure was available.



BACKGROUND

Description of the condition

One of the common features of psychotherapies and professional interventions for people with schizophrenia is to provide support. For example, in an out-patient consultation there will often be time allocated to listening to patients' concerns, giving encouragement, or even arranging basic help with day to day living, such as access to local resources. The key elements of support are to maintain an existing situation, or assist pre-existing coping abilities. Supportive therapy has been described as the treatment model of choice for most patients (Hellerstein 1994), and may be the most commonly practiced intervention in the mental health service.

Description of the intervention

It is, however, difficult to answer the question of exactly what is supportive therapy? A starting point is to look at the literature on supportive psychotherapy, which has been defined as "a dyadic treatment characterised by the use of direct measures to ameliorate symptoms and to maintain, restore or improve self esteem, adaptive skills and ego function" (Pinsker 1991). There may also be a difference in the practice of supportive therapy according to country (Holmes 1995). In the UK, supportive therapy implies a frequency of less than once a week, whereas in America, some practitioners would regard any psychodynamic intervention at a frequency of less than four or five times a week as supportive psychotherapy (Werman 1994).

How the intervention might work

Even though there is no internationally agreed definition, one of the key features is that supportive therapy aims to enhance, rather than challenge, current psychological defence mechanisms. An alternative view is to identify supportive therapy according to the components of the therapy (Misch 2000). For example, one expert proposes seven distinct components; reassurance, explanation, guidance, suggestion, encouragement, affecting change in the environment, and permission for catharsis (Bloch 1996). Bloch also argues that supportive therapy must be a long-term intervention. The difficulty of this proposal is that there is no clear reason why a specific component of therapy should be regarded as supportive. Possible solutions include defining supportive therapy according to outcome, according to the perception of the client or by identifying a feature of therapy that is inherently supportive, regardless of its impact on a client (Barber 2001).

There are many other forms of support that can be given which are distinct from the psychodynamic tradition. For example, there are mental health workers who have the job title of support worker whose role is often to provide practical support such as reminders and transport for other services, or to spend time befriending a client.

Why it is important to do this review

Ultimately, the lack of a widely accepted definition of supportive therapy means we are not able to avoid an arbitrary element in the definition used for this review. Our definition (see Types of interventions) is influenced by the potential usefulness of this review, which could be:

- to give information on the effectiveness of a therapy that is commonly used as the control arm of clinical trials for psychotherapies in schizophrenia;
- 2. to help clinicians with the decision of whether to offer a supportive intervention;
- to help clinicians understand the value of supportive elements of their individual interactions with people who have schizophrenia.

OBJECTIVES

To review the effects of supportive therapy compared with standard care, or other treatments in addition to standard care for people with schizophrenia.

METHODS

Criteria for considering studies for this review

Types of studies

We sought all relevant randomised controlled trials. Where a trial was described as 'double-blind', but it was only implied that the study was randomised, we included these trials in a sensitivity analysis. If there was no substantive difference within primary outcomes (see 'Types of outcome measures') when these 'implied randomisation' studies were added, then we included them in the final analyses. If there was a substantive difference, then we only used clearly randomised trials and the results of the sensitivity analysis were described in the text. We excluded quasi-randomised studies, such as those allocating by using alternate days of the week.

Types of participants

We included people with schizophrenia or schizophrenia-like illnesses using any criteria. We included trials where it was implied that the majority of participants had a severe mental illness that was likely to be schizophrenia. We did not exclude trials due to age, nationality, gender, duration of illness or treatment setting.

Types of interventions

1. Supportive therapy and supportive care

We have used the phrase 'supportive therapy and supportive care' here to indicate that this review covers a wider variety of interventions than supportive psychotherapy alone. However, for the sake of simplicity, we used the term 'supportive therapy' elsewhere in this review. These interventions are provided by a single person with the main purpose of maintaining current functioning or assisting pre-existing coping abilities in people who have a diagnosis of schizophrenia or schizophrenia-like illness. The therapies can be aimed at individuals or groups of people. If the content of the therapy was not sufficiently clear after reading a clinical study, then we included any therapy that had supportive or support in its title. We have included advocacy as a form of supportive therapy. Advocacy is a narrower intervention than other interventions included in the review, but it nevertheless fits our definition as it assists people with their communication and interaction with mental health workers.

A number of common therapies are excluded as they are designed to teach new skills or change pre-existing skills. These include cognitive behavioural therapy(CBT) (Cormac 2004), social skills



training, psycho-education (Pekkala2002), compliance therapy (McIntosh 2006) and problem-solving therapy.

Some therapies or schemes have been excluded as they involve a team approach rather than an individual worker, or because they are designed to alter a person's environment, rather than to help the person cope with their environment. These include family placement (Pharoah 2006), supported employment and supported accommodation (Chilvers 2006) . An exception was to include an intervention if it was clear that it consisted of a practitioner whose main purpose was to help a client cope with their current situation, rather than alter the situation to make it easier for the client.

We have also excluded counselling (unlike the meta-analysis from the National Institute of Clinical Excellence on supportive therapy (NICE 2003)). The main purpose of counselling is to give an opportunity for a client to explore, discover and clarify ways of living (DoH 2001). Counselling employs a wide variety of techniques, and the purpose may be to facilitate a change in someone's life rather than to maintain the current situation. As counselling has such a broad scope, an exception was to include a clinical trial if there was a clear indication that the main purpose of counselling was to provide support rather than facilitate change or give an opportunity to explore personal issues.

2. Standard care

This is defined as the care a person would normally receive had they not been included in the research trial. This would include interventions such as medication, hospitalisation, community psychiatric nursing input and/or day hospital.

3. Other treatments

This would include any other treatment (biological, psychological or social) such as medication, problem-solving therapy, psychoeducation, social skills training, CBT, family therapy or psychodynamic psychotherapy.

Types of outcome measures

We reported outcomes for the short term (up to 12 weeks), medium term (13 to 26 weeks), and long term (more than 26 weeks)

Primary outcomes

1. Global state

1.1 Relapse

2. Service outcomes

2.1 Hospitalisation

5. General functioning

5.1 No clinically important change in general functioning

Secondary outcomes

1. Global state

- 1.2 Time to relapse
- 1.3 No clinically important change in global state
- 1.4 Not any change in global state
- 1.5 Average endpoint global state score
- 1.6 Average change in global state scores

2. Service outcomes

2.2 Time to hospitalisation

3. Mental state

- 3.1 No clinically important improvement
- 3.2 Not any change in general mental state
- 3.3 Average endpoint general mental state score
- 3.4 Average change in general mental state scores
- 3.5 No clinically important change in specific symptoms
- 3.6 Not any change in specific symptoms
- 3.7 Average endpoint specific symptom score
- 3.8 Average change in specific symptom scores

4. Leaving the study early

- 4.1 For specific reasons
- 4.2 For general reasons

5. General functioning

- 5.2 Not any change in general functioning
- 5.3 Average endpoint general functioning score
- 5.4 Average change in general functioning scores
- 5.5 No clinically important change in specific aspects of functioning, such as social or life skills
- 5.6 Not any change in specific aspects of functioning, such as social or life skills
- 5.7 Average endpoint specific aspects of functioning, such as social or life skills
- $5.8\,\mathrm{Average}$ change in specific aspects of functioning, such as social or life skills

6. Satisfaction with treatment

- 6.1 Recipient of care not satisfied with treatment
- 6.2 Recipient of care average satisfaction score
- 6.3 Recipient of care average change in satisfaction scores
- 6.4 Carer not satisfied with treatment
- 6.5 Carer average satisfaction score
- 6.6 Carer average change in satisfaction scores

7. Quality of life

- 7.1 No clinically important change in quality of life
- 7.2 Not any change in quality of life
- 7.3 Average endpoint quality of life score
- 7.4 Average change in quality of life scores
- 7.5 No clinically important change in specific aspects of quality of life
- 7.6 Not any change in specific aspects of quality of life
- 7.7 Average endpoint specific aspects of quality of life
- 7.8 Average change in specific aspects of quality of life

8. Death - suicide and natural causes

9. Behaviour

- 9.1 No clinically important change in general behaviour
- 9.2 Not any change in general behaviour
- 9.3 Average endpoint general behaviour score
- 9.4 Average change in general behaviour scores
- 9.5 No clinically important change in specific aspects of behaviour
- 9.6 Not any change in specific aspects of behaviour
- 9.7 Average endpoint specific aspects of behaviour
- 9.8 Average change in specific aspects of behaviour



10. Adverse effects

- 10.1 No clinically important general adverse effects
- 10.2 Not any general adverse effects
- 10.3 Average endpoint general adverse effect score
- 10.4 Average change in general adverse effect scores
- 10.5 No clinically important change in specific adverse effects
- 10.6 Not any change in specific adverse effects
- 10.7 Average endpoint specific adverse effects
- 10.8 Average change in specific adverse effects

11. Engagement with services

- 11.1 No clinically important engagement
- 11.2 Not any engagement
- 11.3 Average endpoint engagement score
- 11.4 Average change in engagement scores

12. Engagement in structured activities

- 12.1 No clinically important change in engagement in structured activities
- 12.2 Not any change in engagement in structured activities
- 12.3 Average endpoint engagement in structured activities score
- 12.4 Average change in engagement in structured activities scores
- 12.5 No clinically important change in specific activities, such as employment, education or attendance at day centres
- 12.6 Not any change in specific aspects of functioning, such as employment, education or attendance at day centres
- 12.7 Average endpoint specific aspects of functioning, such as employment, education or attendance at day centres
- 12.8 Average change in specific aspects of functioning, such as employment, education or attendance at day centres

13. Insight

- 13.1 Average endpoint insight score
- 13.2 Average endpoint treatment attitude score
- 13.3 Average endpoint adherence score

14. Compliance

- 14.1 Adherence
- 14.2 Poor Compliance

15. Medication

- 15.1 Reduction in dose
- 15.2 Prescribed intramuscular (IM) depot medication

16. Attitude to medication

- 16.1 Average endpoint attitude to medication score
- 16.2 Average endpoint need for treatment

17. Economic outcomes

17.1 Average change in total cost of medical and mental health care 17.2 Total indirect and direct costs.

18. 'Summary of findings' table

We used the GRADEapproach to interpret findings (Schünemann 2008) and used the GRADE profiler to import data from Review Manager (RevMan) to create 'Summary of findings' tables. These tables provide outcome-specific information concerning the overall quality of evidence from each included study in the comparison, the magnitude of effect of the interventions examined, and the sum of available data on all outcomes we rated as

important to patient-care and decision making. We selected the following main outcomes for inclusion in the 'Summary of findings' tables.

- 1. Global state: Relapse
- 2. Service outcomes: Hospitalisation
- 3. Leaving the study early
- 4. Mental state: No clinically important change in general mental state
- General functioning: No clinically important change in general functioning
- 6. Satisfaction with treatment: Recipient of care not satisfied with treatment
- 7. Quality of life: No clinically important change in quality of life

Search methods for identification of studies

Electronic searches

1. Cochrane Schizophrenia Group Trials Register

The Trials Search Co-ordinator searched the Cochrane Schizophrenia Group's Trials Register (28 November 2012)

[((*support* OR *advoc*) in title of REFERENCE) and ((*support* or *individual* or *sociotherapy* or *socioenvir*) in intervention of STUDY)]

The Cochrane Schizophrenia Group's Trials Register is compiled by systematic searches of major databases, handsearches of relevant journals and conference proceedings. For details of databases searched see Group Module.

Searching other resources

1. Reference searching

We inspected the reference lists of all identified studies, including existing reviews, for relevant citations.

2. Personal contact

If necessary we would have contacted the first author of each relevant study for information on unpublished trials.

Data collection and analysis

Selection of studies

For this update review authors NM and KSW independently inspected citations from the new electronic search and identified relevant abstracts. NM and KSW also inspected full articles of the abstracts meeting inclusion criteria. CEA carried out the reliability check of all citations from the new electronic search.

Data extraction and management

1. Extraction

For this update, NM extracted data from included studies. If data were presented only in graphs and figures NM extracted these data whenever possible. If studies were multi-centre, where possible, NM extracted data relevant to each component centre separately.



2. Management

2.1 Forms

We extracted data onto standard, simple forms.

2.2 Scale-derived data

We included continuous data from rating scales only if: a. the psychometric properties of the measuring instrument have been described in a peer-reviewed journal (Marshall 2000); and b. the measuring instrument has not been written or modified by one of the trialists for that particular trial.

Ideally, the measuring instrument should either be i. a self-report or ii. completed by an independent rater or relative (not the therapist). We realise that this is not often reported clearly; if relevant we noted whether or not this is the case in Description of studies.

2.3 Endpoint versus change data

There are advantages of both endpoint and change data. Change data can remove a component of between-person variability from the analysis. On the other hand, calculation of change needs two assessments (baseline and endpoint), which can be difficult in unstable and difficult to measure conditions such as schizophrenia. We decided primarily to use endpoint data, and only use change data if the former were not available. Had we had both change and endpoint data, we would have combined them in the analysis as we used mean differences (MD) rather than standardised mean differences (SMD) throughout (Higgins 2011, Chapter 9.4.5.2).

2.4 Skewed data

Continuous data on clinical and social outcomes are often not normally distributed. To avoid the pitfall of applying parametric tests to non-parametric data, we aimed to apply the following standards to all data before inclusion:

- standard deviations (SDs) and means are reported in the paper or obtainable from the authors;
- when a scale starts from the finite number zero, the SD, when multiplied by two, is less than the mean (as otherwise the mean is unlikely to be an appropriate measure of the centre of the distribution (Altman 1996));
- if a scale started from a positive value (such as the Positive and Negative Syndrome Scale (PANSS, (Kay 1986), which can have values from 30 to 210), we modified the calculation described above to take the scale starting point into account. In these cases skew is present if 2 SD > (S-S min), where S is the mean score and S min is the minimum score.

Endpoint scores on scales often have a finite start and end point and these rules can be applied. We entered skewed endpoint data from studies of fewer than 200 participants as other data within the data and analyses section rather than into a statistical analysis. Skewed data pose less of a problem when looking at mean if the sample size is large; we entered such endpoint data into syntheses.

When continuous data are presented on a scale that includes a possibility of negative values (such as change data), it is difficult to tell whether data are skewed or not, we entered skewed change data into analyses regardless of size of study.

2.5 Common measure

To facilitate comparison between trials, we intended to convert variables that can be reported in different metrics, such as days in hospital (mean days per year, per week or per month) to a common metric (e.g. mean days per month).

2.6 Conversion of continuous to binary

Where possible, we made efforts to convert outcome measures to dichotomous data. This can be done by identifying cut-off points on rating scales and dividing participants accordingly into 'clinically improved' or 'not clinically improved'. It is generally assumed that if there is a 50% reduction in a scale-derived score such as the Brief Psychiatric Rating Scale (BPRS, Overall 1962) or the Positive and Negative Syndrome Scale (PANSS, Kay 1986), this could be considered as a clinically significant response (Leucht 2005; Leucht 2005a). If data based on these thresholds were not available, we used the primary cut-off presented by the original authors.

2.7 Direction of graphs

Where possible, we entered data in such a way that the area to the left of the line of no effect indicated a favourable outcome for supportive therapy.

Assessment of risk of bias in included studies

For this update, NM and KSW worked independently by using criteria described in the *Cochrane Handbook for Systematic Reviews* of *Interventions* (Higgins 2011) to assess trial quality. This new set of criteria is based on evidence of associations between overestimate of effect and high risk of bias of the article such as sequence generation, allocation concealment, blinding, incomplete outcome data and selective reporting.

Where inadequate details of randomisation and other characteristics of trials were provided, we contacted authors of the studies in order to obtain additional information.

We have noted the level of risk of bias in both the text of the review and in the 'Summary of findings' tables.

Measures of treatment effect

1. Binary data

For binary outcomes, we calculated a standard estimation of the risk ratio (RR) and its 95% confidence interval (CI). It has been shown that RR is more intuitive (Boissel 1999) than odds ratios and that odds ratios tend to be interpreted as RR by clinicians (Deeks 2000). The Number Needed to Treat/Harm (NNT/H) statistic with its confidence intervals is intuitively attractive to clinicians but is problematic both in its accurate calculation in meta-analyses and interpretation (Hutton 2009). For binary data presented in the 'Summary of findings' tables, where possible, we calculated illustrative comparative risks.

2. Continuous data

For continuous outcomes, we estimated mean difference (MD) between groups. We would prefer not to calculate effect size measures (standardised mean difference (SMD)). However, if scales of very considerable similarity had been used, we would have presumed there was a small difference in measurement, and we would have calculated effect size and transformed the effect back to the units of one or more of the specific instruments.



Unit of analysis issues

1. Cluster trials

Studies increasingly employ 'cluster randomisation' (such as randomisation by clinician or practice), but analysis and pooling of clustered data poses problems. Authors often fail to account for intra-class correlation in clustered studies, leading to a 'unit of analysis' error (Divine 1992) whereby P values are spuriously low, confidence intervals unduly narrow and statistical significance overestimated. This causes type I errors (Bland 1997; Gulliford 1999).

We did not include any cluster-randomised trials in this version of the review. If we had found trials where clustering was not accounted for in primary studies, we would have presented data in a table, with a (*) symbol to indicate the presence of a probable unit of analysis error. In subsequent versions of this review, if we include cluster-randomised trials, we will seek to contact first authors of studies to obtain intra-class correlation coefficients (ICCs) for their clustered data and to adjust for this by using accepted methods (Gulliford 1999).

If we had included trials where clustering had been incorporated into the analysis of primary studies, we would have presented these data as if from a non-cluster randomised study, but adjusted for the clustering effect.

We have sought statistical advice and have been advised that the binary data as presented in a report should be divided by a 'design effect'. This is calculated using the mean number of participants per cluster (m) and the ICC [Design effect = 1+(m-1)*ICC] (Donner 2002). If the ICC is not reported it will be assumed to be 0.1 (Ukoumunne 1999).

If cluster studies have been appropriately analysed taking into account ICCs and relevant data documented in the report, synthesis with other studies would be possible using the generic inverse variance technique.

2. Cross-over trials

A major concern of cross-over trials is the carry-over effect. It occurs if an effect (e.g. pharmacological, physiological or psychological) of the treatment in the first phase is carried over to the second phase. As a consequence, on entry to the second phase the participants can differ systematically from their initial state despite a wash-out phase. For the same reason, cross-over trials are not appropriate if the condition of interest is unstable (Elbourne 2002). As both effects are very likely in severe mental illness, had we included such trials, we planned to use only the data of the first phase of cross-over studies.

3. Studies with multiple treatment groups

Where a study involves more than two treatment arms, if relevant, we presented the additional treatment arms in comparisons. If data were binary, we simply added these and combined within the two-by-two table. If data were continuous, we combined data following the formula in section 7.7.3.8 (Combining groups) of the *Cochrane Handbook for Systemic reviews of Interventions* (Higgins 2011). Where the additional treatment arms were not relevant, we did not reproduce these data.

Dealing with missing data

1. Overall loss of credibility

At some degree of loss of follow-up, data must lose credibility (Xia 2009). We chose that, for any particular outcome, should more than 50% of data be unaccounted for, we would not reproduce these data or use them within analyses. If, however, more than 50% of those in one arm of a study were lost, but the total loss was less than 50%, we addressed this within the 'Summary of findings' tables by down-rating quality. Finally, we also downgraded quality within the 'Summary of findings' tables where loss was 25% to 50% in total.

2. Binary

In the case where attrition for a binary outcome was between 0% and 50% and where these data were not clearly described, we presented data on a 'once-randomised-always-analyse' basis (an intention-to-treat analysis). Those leaving the study early were all assumed to have the same rates of negative outcome as those who completed, with the exception of the outcome of death and adverse effects. For these outcomes, the rate of those who stayed in the study - in that particular arm of the trial - were used for those who did not. We undertook a sensitivity analysis to test how prone the primary outcomes were to change when 'completer' data only were compared to the intention-to-treat analysis using the above assumptions.

3. Continuous

3.1 Attrition

In the case where attrition for a continuous outcome is between 0% and 50% and completer-only data were reported, we reproduced these.

3.2 Standard deviations

If standard deviations (SDs) were not reported, we first tried to obtain the missing values from the authors. If not available, where there are missing measures of variance for continuous data, but an exact standard error (SE) and confidence intervals available for group means, and either a P value or T value available for differences in mean, we can calculate them according to the rules described in the Handbook (Higgins 2011): When only the SE is reported, SDs are calculated by the formula SD = SE * square root (n). Chapters 7.7.3 and 16.1.3 of the Handbook (Higgins 2011) present detailed formulae for estimating SDs from P values, T or F values, confidence intervals, ranges or other statistics. If these formulae do not apply, we can calculate the SDs according to a validated imputation method, which is based on the SDs of the other included studies (Furukawa 2006). Although some of these imputation strategies can introduce error, the alternative would be to exclude a given study's outcome and thus to lose information. We nevertheless examined the validity of the imputations in a sensitivity analysis excluding imputed values.

3.3 Last observation carried forward

We anticipated that in some studies the method of last observation carried forward (LOCF) would be employed within the study report. As with all methods of imputation to deal with missing data, LOCF introduces uncertainty about the reliability of the results (Leucht 2007). Therefore, where LOCF data have been used in the trial, if less than 50% of the data have been assumed, we reproduced these data and indicated that they are the product of LOCF assumptions.



Assessment of heterogeneity

1. Clinical heterogeneity

We considered all included studies initially, without seeing comparison data, to judge clinical heterogeneity. We simply inspected all studies for clearly outlying people or situations which we had not predicted would arise. When such situations or participant groups arose, we fully discussed these.

2. Methodological heterogeneity

We considered all included studies initially, without seeing comparison data, to judge methodological heterogeneity. We simply inspected all studies for clearly outlying methods which we had not predicted would arise. When such methodological outliers arose, we fully discussed these.

3. Statistical heterogeneity

3.1 Visual inspection

We visually inspected graphs to investigate the possibility of statistical heterogeneity.

3.2 Employing the I² statistic

We investigated heterogeneity between studies by considering the I² method alongside the Chi² P value. The I² provides an estimate of the percentage of inconsistency thought to be due to chance (Higgins 2003). The importance of the observed value of I² depends on i. magnitude and direction of effects and ii. strength of evidence for heterogeneity (e.g. P value from Chi² test, or a confidence interval for I2). An I2 estimate greater than or equal to around 50% accompanied by a statistically significant Chi² statistic was interpreted as evidence of substantial levels of heterogeneity (Higgins 2011). When substantial levels of heterogeneity were found in the primary outcome, we explored reasons for heterogeneity (Subgroup analysis and investigation of heterogeneity). Where the I² estimate was equal to or greater than 75%, we interpreted this as indicating the presence of high levels of heterogeneity (Higgins 2003). We did not summate data if inconsistency was high, but presented the data separately and investigated reasons for heterogeneity. If data were heterogeneous we used a random-effects model.

Assessment of reporting biases

Reporting biases arise when the dissemination of research findings is influenced by the nature and direction of results (Egger 1997). These are described in Section 10 of the *Handbook* (Higgins 2011). We are aware that funnel plots may be useful in investigating reporting biases but are of limited power to detect small-study effects. We did not use funnel plots for outcomes where there were 10 or fewer studies, or where all studies were of similar sizes. In other cases, where funnel plots are possible, we sought statistical advice in their interpretation.

Data synthesis

We understand that there is no closed argument for preference for use of fixed-effect or random-effects models. The random-effects method incorporates an assumption that the different studies are estimating different, yet related, intervention effects. This often seems to be true to us and the random-effects model takes into account differences between studies even if there is no statistically

significant heterogeneity. There is, however, a disadvantage to the random-effects model: it puts added weight onto small studies which often are the most biased ones. Depending on the direction of effect, these studies can either inflate or deflate the effect size. We chose the fixed-effect model for all analyses. The reader is, however, able to choose to inspect the data using the random-effects model.

Subgroup analysis and investigation of heterogeneity

1. Subgroup analyses - only primary outcomes

1.1 Clinical state, stage or problem

We proposed to undertake this review and provide an overview of the effects of supportive therapy for people with schizophrenia in general. In addition, however, we tried to report data on subgroups of people in the same clinical state, stage and with similar problems. However, there were not enough included studies in the comparisons to undertake subgroup analyses.

2. Investigation of heterogeneity

If inconsistency was high, we have reported this. First, we investigated whether data had been entered correctly. Second, if data were correct, we visually inspected the graph and successively removed outlying studies to see if homogeneity was restored. For this review we decided that should this occur with data contributing to the summary finding of no more than around 10% of the total weighting, we would present these data. If not, then we did not pool data and discussed issues. We know of no supporting research for this 10% cut-off, but we use prediction intervals as an alternative to this unsatisfactory state.

When unanticipated clinical or methodological heterogeneity was obvious, we simply stated hypotheses regarding these for future reviews or versions of this review. We do not anticipate undertaking analyses relating to these.

Sensitivity analysis

We applied all sensitivity analyses to the primary outcomes of this review.

1. Implication of randomisation

We aimed to include trials in a sensitivity analysis if they were described in some way so as to imply randomisation. For the primary outcomes, we included these studies and if there was no substantive difference when the implied randomised studies were added to those with better description of randomisation, then we entered all data from these studies.

2. Assumptions for lost binary data

Where assumptions had to be made regarding people lost to followup (see Dealing with missing data), we compared the findings of the primary outcomes when we used our assumption/s and when we used data only from people who completed the study to that point. If there was a substantial difference, we reported results and discussed them, but continued to employ our assumption.

Where assumptions had to be made regarding missing SDs data (see Dealing with missing data), we compared the findings of the primary outcomes when we used our assumption/s and when we used data only from people who completed the study to that point. A sensitivity analysis was undertaken to test how prone results



were to change when completer-only data only were compared to the imputed data using the above assumption. If there was a substantial difference, we reported results and discussed them, but continued to employ our assumption

3. Risk of bias

We analysed the effects of excluding trials that were judged to be at high risk of bias across one or more of the domains of randomisation (implied as randomised with no further details available): allocation concealment, blinding and outcome reporting for the meta-analysis of the primary outcome. If the exclusion of trials at high risk of bias did not substantially alter the direction of effect or the precision of the effect estimates, then we included data from these trials in the analysis.

4. Imputed values

We planned to undertake a sensitivity analysis to assess the effects of including data from trials if we had used imputed values for ICC in calculating the design effect in cluster-randomised trials, however, no cluster-randomised trials were included in this version of the review.

If we noted substantial differences in the direction or precision of effect estimates in any of the sensitivity analyses listed above, we

did not pool data from the excluded trials with the other trials contributing to the outcome, but presented them separately.

5. Fixed-effect and random-effects

We synthesised data using a fixed-effect model, unless data were heterogenous, in which case we used a random-effects model (see Assessment of heterogeneity).

RESULTS

Description of studies

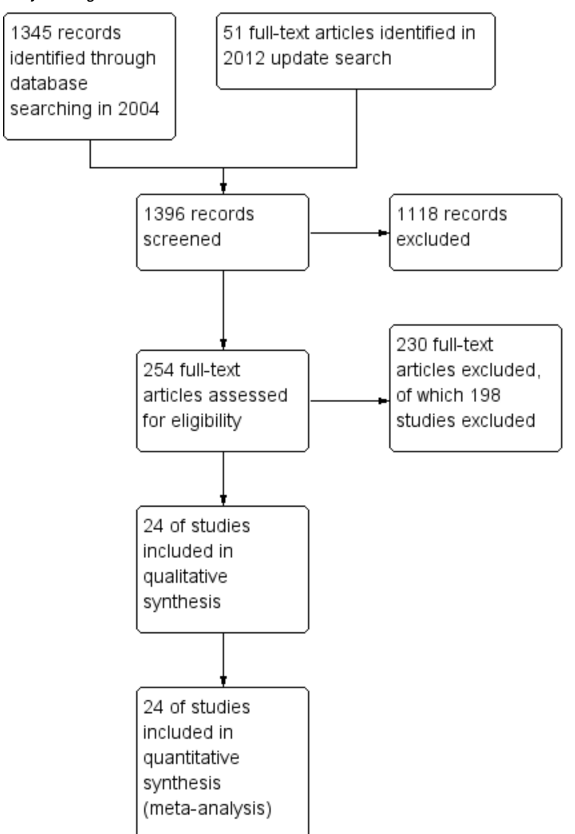
Please see Characteristics of included studies, Characteristics of excluded studies, Characteristics of studies awaiting classification, and Characteristics of ongoing studies.

Results of the search

In total, we identified 1396 references from the searches, many of which could be excluded on the basis of their abstracts. We selected 227 references in the previous search and 51 from the update search and obtained full-text papers for assessment. We included an additional four studies (Davidson 2004; Klein 1998; Penn 2009; Uzenoff 2007); the review now includes 24 studies with 56 references (see Figure 1).



Figure 1. Study flow diagram.





Included studies

1. Method

Due to the nature of the interventions, none of the 24 included studies were able to use a double-blind technique. Some (11/24), however, endeavoured to reduce bias by having some or all outcomes rated by people blind to allocation (Durham 2003; Eckman 1992; Haddock 1999; Kemp 1996; Levine 1998; Lewis 2002b; Sensky 2000b; Spaulding 1999; Stanton 1984; Tarrier 1998; Turkington 2000). Two studies attempted to keep therapists blind to the specific study hypothesis (Coyle 1988; Stanton 1984). Some studies reported outcomes immediately after therapy, whilst others reported outcomes after a follow-up period without therapy.

2. Duration

The overall duration of the trials varied between 10 weeks and three years. Three studies (Levine 1998; Lewis 2002b; Penn 2009) were short term (up to 12 weeks). Five studies (Coyle 1988; Klein 1998; Pinto 1999; Turkington 2000; Uzenoff 2007) were medium term (13 to 26 weeks), and the remaining 16 studies were long term (more than 26 weeks).

3. Participants

Nineteen included studies had less than 100 participants; the range of the number of participants was 12 to 315. A total of 2126 people participated in the 24 trials, most of whom had a diagnosis of schizophrenia. Ten studies included only people with schizophrenia, however two of these (Coyle 1988; Malm 1982) did not describe criteria clearly. Sixteen trials included people with other psychotic illnesses (schizoaffective disorder, delusional disorder and other psychoses). Of these, one study also included people with bipolar disorder (O'Donnell 1999) and one included severely disturbed psychiatric patients (Dincin 1982). Twentythree studies employed operational criteria for diagnoses (DSM-II, DSM-III, DSM-III-R, DSM-IV, ICD-10, RDC, WHO Discrimination Criteria), however, one of these trials Coyle 1988 did not report how the diagnoses were reached. Malm 1982 described the diagnostic criteria used, but these were not recognised operational criteria. Twenty trials included male and female participants, two studies (Eckman 1992; Wirshing 1991) included men only, and the remaining two studies (Levine 1998; Uzenoff 2007) did not report the sex of participants. Ages of participants ranged from 16 to 72 years.

4. Setting

Eleven trials took place in outpatient settings. Three trials were conducted in inpatient settings (Haddock 1999; Spaulding 1999; Stanton 1984), seven were in a mixture of inpatient and outpatient settings (Durham 2003; Lewis 2002b; Malm 1982; O'Donnell 1999; Pinto 1999; Turkington 2000; Wirshing 1991), and the rest did not report on setting.

Thirteen studies were carried out in the USA (Coyle 1988; Davidson 2004; Dincin 1982; Eckman 1992; Falloon 1982; Hogarty 1997-study 1; Hogarty 1997-study 2; Klein 1998; Penn 2009; Spaulding 1999; Stanton 1984; Uzenoff 2007; Wirshing 1991); seven in the UK (Durham 2003, Haddock 1999; Kemp 1996; Lewis 2002b; Sensky 2000b; Tarrier 1998; Turkington 2000); one in Australia (O'Donnell 1999); one in Israel (Levine 1998); one in Italy (Pinto 1999); and one in Sweden (Malm 1982).

5. Interventions

5.1 Experimental intervention

All studies used supportive therapy in addition to standard care (including antipsychotic medication). There were variations between studies with regard to frequency and duration of therapy sessions. Most studies used twice-weekly, weekly and fortnightly sessions, although some studies did not report on the frequency or duration of intervention (Durham 2003; Haddock 1999; O'Donnell 1999; Pinto 1999). The duration of therapy varied between five weeks and three years. Fifteen studies investigated individual treatment, and six studies delivered supportive therapy in a group format (Dincin 1982; Eckman 1992; Levine 1998; Malm 1982; Spaulding 1999; Wirshing 1991). Seven studies reported the use of therapy manuals or protocols (Durham 2003; Hogarty 1997study 1; Hogarty 1997-study 2; Lewis 2002b; Malm 1982; Penn 2009; Spaulding 1999). Eight studies reported that therapists were supervised (Davidson 2004; Haddock 1999; Hogarty 1997-study 1; Hogarty 1997-study 2; Lewis 2002b; Malm 1982; Penn 2009; Sensky 2000b), and six that audiotapes of sessions were monitored for quality (Haddock 1999; Lewis 2002b; Penn 2009; Sensky 2000b; Turkington 2000; Uzenoff 2007). Six of the studies reported that the same therapists delivered experimental and control interventions, and three studies reported using different therapists for different therapeutic interventions (Durham 2003; O'Donnell 1999; Stanton 1984). Therapists delivering the supportive intervention were trained in a different therapeutic modality, such as cognitive behavioural therapy (CBT) or family therapy, in four studies (Haddock 1999; Lewis 2002b; Penn 2009; Sensky 2000b).

5.2 Control intervention

Five studies compared supportive therapy with standard treatment alone (Coyle 1988; Davidson 2004; Durham 2003; Lewis 2002b; Tarrier 1998), the remaining trials used various other psychosocial interventions for comparison. Twelve studies compared supportive therapy with CBT (Durham 2003; Haddock 1999; Hogarty 1997study 1; Hogarty 1997-study 2; Kemp 1996; Levine 1998; Lewis 2002b; Pinto 1999; Sensky 2000b; Spaulding 1999; Tarrier 1998; Turkington 2000). Two studies used family therapy as a comparison (Falloon 1982; Hogarty 1997-study 1). Skills training was investigated in three studies (Coyle 1988; Eckman 1992; Wirshing 1991); other comparisons were personal therapy plus family therapy (Hogarty 1997-study 1), psychoeducation (Coyle 1988; Uzenoff 2007), milieu rehabilitation programme (Dincin 1982) and insight-oriented psychotherapy (Stanton 1984). One study investigated supportive therapy combined with client-focused case management in comparison with client-focused case management (O'Donnell 1999). One trial compared supportive therapy with intensive case management in comparison with intensive case management (Klein 1998). Finally, one trial investigated the effect of adding supportive therapy to a combination of social skills training and medication (Malm 1982). Fourteen of the studies attempted to match experimental and control psychosocial interventions for the amount of therapist contact (Eckman 1992; Falloon 1982; Haddock 1999; Kemp 1996; Levine 1998; Lewis 2002b; Penn 2009; Pinto 1999; Sensky 2000b; Spaulding 1999; Tarrier 1998; Turkington 2000; Uzenoff 2007; Wirshing 1991). In contrast, four studies took the approach that different interventions by their nature involve different amounts of therapist contact (Dincin 1982; Hogarty 1997-study 1; Hogarty 1997-study 2; Stanton 1984). The other studies did not report on this matter. Davidson 2004 gave all participants a \$28 stipend whether they received supportive care or



not to control for possible effects of receiving funds to take part in social activities.

6.Outcomes

Listed below are the outcomes for which we could obtain usable data, followed by a summary of data that could not be used in the meta-analysis.

6.2 Outcome scales

6.2.1 Mental state

i. Positive and Negative Syndrome Scale (PANSS) (Kay 1987)

This scale assesses the severity of psychotic symptomatology in general. It consists of three subscales, positive symptoms, negative symptoms and general psychopathology, and a total score. This scale was used by Durham 2003; Levine 1998; Lewis 2002b; Penn 2009 and Uzenoff 2007.

ii. Brief Psychiatric Rating Scale (BPRS) (Lukoff 1996)

This is a semi-structured interview, which assesses the major psychiatric symptoms. It is a 16-item scale, and each item is scored on a seven-point scale, ranging from 'not present' to 'extremely severe'. The range of possible scores is 24 to 168, and high scores indicate more severe symptoms. Data from the BPRS is reported by Haddock 1999; Kemp 1996; Penn 2009; and Pinto 1999. Kemp 1996 reported data for the full version of the BPRS immediately after treatment. They also used an abridged version of the scale, which contained seven of the 16 items, including psychotic symptoms, negative symptoms and depression. Results for the abridged version were reported immediately after treatment and at six-month, 12-month and 18-month follow-up.

iii. Schedule for Assessment of Negative Symptoms (SANS) (Andreasen 1983)

This scale assesses the negative symptoms of schizophrenia. A six-point scale is used to rate alogia, affective blunting, avolition-apathy, anhedonia-apathy, anhedonia-asociality and attention impairment. Higher scores indicate more severe symptoms. This scale was used by Pinto 1999 and Sensky 2000b.

iv. Schedule for Assessment of Positive Symptoms (SAPS) (Andreasen 1984)

This scale selectively assesses the positive symptoms of psychosis and the higher the score, the more severe the symptoms. Pinto 1999 used this scale.

v. Psychotic Symptoms Rating Scales (PSYRATS) (Haddock 1999b) This consists of two scales, which assess delusional beliefs and auditory hallucinations. There are 11 items in the auditory hallucinations scale, including frequency, duration, level of distress, controllability, loudness, location and beliefs about origin of voices. The delusional beliefs scale has six items, including preoccupation, intensity of distress, conviction and disruption. Each item is rated on a five-point scale. The PSYRATS were used by Lewis 2002b and Penn 2009.

vi. Belief about Voices Questionnaire-Revised (BAVQ-R) (Chadwick 2000)

This scale is a 35-item measure of beliefs about auditory hallucinations and the emotional and behavioural reactions to them. There are five BAVQ-R subscales: malevolence, benevolence, resistance, engagement, and omnipotence. Each item is rated on a

four-point scale; a higher score indicates more belief in their voices. This scale was used by Penn 2009.

vii. Comprehensive Psychiatric Rating Scale (CPRS) (Montgomery 1978)

This is a general psychiatric rating scale, which was used by Sensky 2000b and Malm 1982. A high score indicates severe symptoms. Malm 1982 used the schizophrenia subscale, which consists of 45 items and a global rating of degree of illness. The paper reported useable data for only two of the 45 items on this scale.

viii. Beck Depression Inventory II (BDI-II) (Beck 1996)

This scales measures the severity of depression. It has 21 self-reported items measured on a four point scale. A higher score indicates more severe depression. This scale was used by Penn 2009.

ix. Calgary Depression Rating Scale (CDRS) (Addington 1993)

This is a nine-item structured interview scale developed specifically for assessing depression in individuals with schizophrenia. The CDRS assesses depression as separate from overlapping negative or extrapyramidal symptoms. Higher scores indicate greater depression. This scale was used by Uzenoff 2007.

x. Center for Epidemiological Studies—Depression Scale (CES-D) (Radloff 1977)

This is a 20-item scale to measure depressive symptomology. The possible range of scores is zero to 60 and higher scores indicate more symptoms. This scale was used by Davidson 2004.

xi. Montgomery-Asberg Depression Rating Scale (MADRS) (Montgomery 1979)

This scale was developed using a 65-item psychopathology scale to identify the 17 most commonly occurring symptoms in primary depressive illness. The maximum score is 30, and a higher score indicates more severe psychopathology. The scale was used by Sensky 2000b.

6.1.2 General functioning

i. Global Assessment of Functioning Scale (GAF) (APA 1987) The GAF disability scale is taken from the DSM-III-R (APA 1987). The scale has 10 defined anchor points relating to social competence, and scores range from zero to 100. A higher score indicates better functioning. Durham 2003; Kemp 1996; and Klein 1998 report data from this scale. Davidson 2004 uses the GAF-M, which is a modified version.

ii. Global Assessment Scale (GAS) (Endicott 1976)

This scale provides a rating between zero and 100, which reflects overall degree of psychological or psychiatric health. A high score indicates better health. This scale was used by Durham 2003.

iii. Katz Adjustment Scales (KAS) (Katz 1963)

This collection of scales measures social adjustment and behaviour. It was used by Malm 1982. It consists of items which can be grouped into subscales including 'performance of socially expected activities' (15 items) and 'free-time activities' (23 items), the two subscales chosen by Malm 1982. In this study, the items were also grouped into syndromes, entitled 'offensive behaviour' (two items) and 'withdrawal' (six items).

iv. Social Functioning Scale (SFS) (Birchwood 1990)



The SFS is a 79-item self-report questionnaire to both the patients and the caregiver, about performance in seven areas: Social Engagement (SE), Interpersonal Communication (IC), Recreational Activities (RA), Social Activities (SA), Independence Competence (INC), Independence Performance (IP) and Occupational Activity (OA). The purpose of the scale is to provide an evaluation of strengths and weaknesses of patient functioning, and it may reveal aims for therapeutic intervention.

6.1.3 Behaviour

i. Social Behaviour Adjustment Scale (SBAS) (Platt 1980)

Falloon 1982 was the only study to use this scale. Scoring is based on a structured interview with a key informant from the person's family. Areas covered include household tasks, spare time/leisure activity, work/study, decision making, friendliness/affection, everyday conversation, relationships outside the family, behavioural disturbance, social and interpersonal role performance, and adverse effects on the family and other people in the community. Each item is scored on a scale from zero to two, and higher scores indicate greater impairment or dissatisfaction.

6.1.4 Insight

i. Beck Cognitive Insight Scale - (BCIS) composite (Beck 2004)
The BCIS is a 15-item self-report measure composed of two subscales: self-reflectiveness and self-certainty. Penn 2009 used a composite index of insight, computed from the BCIS. Higher scores reflect greater cognitive insight.

ii. Insight and Treatment Attitudes Questionnaire (ITAQ) (McEvoy 1989)

This scale assesses an individual's recognition of illness and need for treatment and includes 11 questions. Reponses are rated as follows: 2 = good insight, 1 = partial insight, and 0 = no insight, which are summed to provide a total insight score. Higher scores correspond to better insight. Uzenoff 2007 reported data on this scale.

iii. Schedule for Assessment of Insight (SAI) (David 1990)

The schedule is a semi-structured interview with three components; treatment compliance, awareness of illness and ability to relabel psychotic symptoms correctly. The range of possible scores is zero to 14, but scores are expressed as a percentage of maximum insight. Low scores therefore indicate poor insight. The expanded version of the SAI was used by Kemp 1996.

6.1.5 Compliance

i. Rating of Medication Influences (ROMI) scale (Weiden 1994)
The ROMI is a comprehensive measure of attitudes regarding adherence to antipsychotic medications used by Uzenoff 2007.
The measure is divided into two subscales, 'Reasons for Adherence' (nine items) and 'Reasons for Nonadherence' (10 items), each containing attitudinal and behavioural factors thought to influence adherence behaviour. Items are rated on a one through three scale, where 1 = no influence, 2 = mild influence, 3 = strong influence, and 96 = not applicable. Higher scores on the 'Reasons for Adherence' subscale correspond to an endorsement of factors influencing the individual to take his/her medication, whereas higher scores on the 'Reasons for Nonadherence' subscale correspond to an endorsement of factors influencing the individual to stop taking his/her medication.

6.1.6 Attitude to medication

i. Attitudes to Medication Questionnaire (AMQ) (Hayward 1995) The AMQ is a semi-structured interview designed to assess attitudes to drug treatment. It was used by Kemp 1996. A high score indicates a helpful attitude to treatment.

ii. Drug Attitudes Inventory (DAI) (Hogan 1983)

The DAI is a self-report scale used to assess attitude to medication. It has been shown to be highly predictive of compliance. A higher score indicates a better attitude to treatment. This scale was used by Kemp 1996.

6.1.7 Quality of life

i. Global Health Questionnaire - GHQ (Goldberg 1972)

This 60-item scale measures the severity of nonpsychotic psychiatric symptomatology. It was used by Davidson 2004. It assesses the respondent's current state and asks if that differs from his or her usual state and produces an overall score that can be compared with a prescribed cut-off score. All items have a four-point Likert scoring (0-1-2-3) system that ranges from a 'better/ healthier than normal' option, through a 'same as usual' and a 'worse/more than usual' to a 'much worse/more than usual' option. The higher the score, the more severe the condition.

ii. Quality of Life Scale - QLS (Heinrichs 1984)

This 21-item scale is based on a semi-structured interview providing information on symptoms and functioning during the preceding four weeks. There are seven severity steps (zero to six; six being adequately functioning and zero being deficient). Four item categories have been identified by factor analysis i) interpersonal relationships (seven items), ii) instrumental role (four items), iii) intrapsychic function (seven items) and iv) common place objects and activities. Higher scores indicate better quality of life. This scale was used by Uzenoff 2007.

iii. Rosenberg Self-Esteem Scale (RSES) (Rosenberg 1965)

The scale measures state self-esteem by asking the respondents to reflect on their current feelings, with a 10-item Likert-type scale with items answered on a four-point scale — from strongly agree to strongly disagree. Five of the items have positively worded statements and five have negatively worded ones. The scale ranges from zero to 30. Scores between 15 and 25 are within normal range; scores below 15 suggest low self-esteem. This scale was used by Davidson 2004 and Penn 2009.

iv. Well-Being scale- WBS (Ryff 1989)

This is a scale to measure positive aspects of psychological functioning along six dimensions in which respondents rate statements on a scale of one to six, with one indicating strong disagreement and six indicating strong agreement: independence and self-determination (autonomy); having satisfying, high quality relationships (positive relations with others); the ability to manage one's life (environmental mastery); being open to new experiences (personal growth); believing that one's life is meaningful (purpose in life); and a positive attitude towards oneself and one's past life (self-acceptance). A high score indicates that the respondent has a mastery of that area in his or her life. Conversely, a low score shows that the respondent struggles to feel comfortable with that particular concept. Davidson 2004 reported data for this scale.

v. Lehman's Quality-of-Life (QOL) Questionnaire (Lehman 1988)



Lehman's QOL scale uses a response range of one (terrible) to seven (delighted). Klein 1998 used only those domains that had direct applicability - namely, living situation, income, family, friends, social interaction, activities of daily living, and health. They also added a section on drug use, as well as a general list of activities, the response scale was one (daily) to five (never).

6.2 Missing outcomes

No studies provided data regarding clinically important change in general functioning, one of our primary outcomes of interest. No studies reported data on time to relapse or hospitalisation, clinically important or any changes in general behaviour scores, adverse events, or in any engagement in structured activities. Economic data, as well as satisfaction with treatment were seldom reported.

Excluded studies

We excluded 198 reports - most because the reported interventions did not meet our criteria for supportive therapy. In several studies, all participants received supportive therapy, without a comparison group relevant to this review. We excluded some studies because the participants did not have schizophrenia or schizophrenia-like illnesses. Some studies were not randomised controlled trials,

and we therefore excluded these. We excluded one study due to inadequate allocation concealment (Slavinsky 1982). Eight studies were excluded as they did not report usable data (Castelein 2008; Castilla-Puentes 2002; Keshavan 2011; Klingberg 2010; Meister 2010; Nelson 2007; Smith 1999; Telles 1995). Details of excluded studies are detailed in the Characteristics of excluded studies.

Awaiting assessment studies

Three studies are awaiting assessment. Bechdolf 2012, and Blankertz 1997 are awaiting translation from German and Falloon 1983 is awaiting assessment until we obtain a full-text paper.

Ongoing studies

Sixteen studies seem to be ongoing. We are obtaining more details to ensure that this is indeed the case. These studies are using a variety of interventions, including behavioural treatment, CBT, cognitive enhancement therapy, body orientated therapy, with supportive therapies and counselling as controls.

Risk of bias in included studies

See also 'Risk of bias' tables in Characteristics of included studies and Figure 2 and Figure 3.

Figure 2.

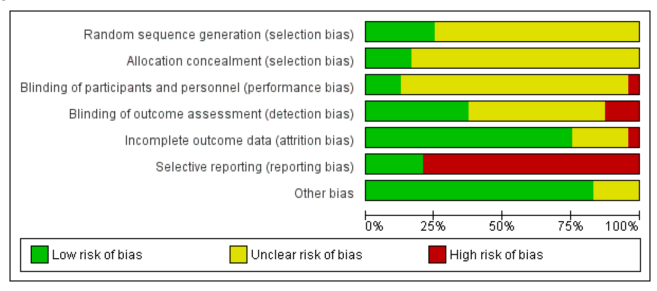


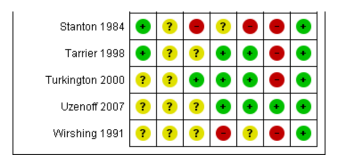


Figure 3.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Coyle 1988	?	?	?	?	•	•	?
Davidson 2004	?	?	?	?	?	•	•
Dincin 1982	?	?	?	?	?	•	•
Durham 2003	•	•	?	?	•	•	•
Eckman 1992	?	?	?	?	?	•	•
Falloon 1982	?	?	?	•	•	•	•
Haddock 1999	?	?	?	•	•	•	?
Hogarty 1997-study 1	?	?	?	•	•	•	•
Hogarty 1997-study 2	?	?	?	•	•	•	•
Kemp 1996	•	?	?	?	•	•	•
Klein 1998	?	?	?	?	•	•	•
Levine 1998	?	?	?	•	?	•	?
Lewis 2002b	•	•	•	•	•	•	•
Malm 1982	?	?	?	?	•	•	•
O'Donnell 1999	?	?	?	?	•	•	•
Penn 2009	•	•	?	•	•	•	•
Pinto 1999	?	?	?	?	•	•	?
Sensky 2000b	?	•	?	•	•	•	•
Spaulding 1999	?	?	•	?	•	•	•
Stanton 1984	•	?		?		•	•



Figure 3. (Continued)



Allocation

All studies were randomised trials. However, most studies did not provide explicit details of the method of randomisation, and were rated as unclear risk of bias. Six studies were rated low risk of bias. Kemp 1996 reports the use of a table of random numbers. Penn 2009 stratified according to gender, and stratification was also undertaken by Lewis 2002b with the following variables: first or second admission; inpatient or day patient admission; male or female; with the first-episode cases further stratified for duration of symptoms of more or less than six months. Tarrier 1998 used block randomisation, with a block size of nine, stratified by severity of symptoms and sex. Durham 2003 used permuted blocking. Spaulding 1999 matched pairs and assigned each paired participant to one of two groups. The groups were then were randomly assigned to either the treatment or the control intervention. Coyle 1988 describes stratifying participants according to age, sex and race prior to randomisation, but no further details were reported and was rated as unclear.

With regard to allocation concealment, most studies did not report any concealment approach, and were rated as unclear risk of bias. Five studies described the method of allocation concealment, four of which we rated as low risk. Durham 2003 provided a clear description of the randomisation procedure, which was administered centrally by a non-clinical project coordinator, and a sealed envelope technique was used. Sensky 2000b reported that randomisation was done by members of the research team not involved with assessment or treatment. Lewis 2002b reported that independent, concealed randomisation with minimisation was performed by trial administrators. Penn 2009 reported randomisation by a blinded researcher, using random numbers generated by computer. Tarrier 1998 reported that randomisation was carried out independently by a third party, using sealed envelopes, but not whether the envelopes were opaque and so we rated it as having an unclear risk of bias.

Blinding

None of the studies were able to use a double-blind technique due to the nature of the interventions, since it would not be possible to have therapists and participants blind to treatment allocation. Turkington 2000, Lewis 2002b, and Spaulding 1999 were the only studies to report appropriate personnel blinding and were rated as low risk of bias. In Stanton 1984, ward and project staff were not blinded, and so was rated as high risk of bias. Blinding of personnel for the remaining trials was not adequately reported and these were rated as unclear risk of bias.

Nine studies stated that outcome raters were blinded to the treatment group (Falloon 1982; Haddock 1999; Levine 1998; Lewis 2002b; Penn 2009; Sensky 2000b; Tarrier 1998; Turkington 2000; Uzenoff 2007) and were rated as low risk of bias for outcome assessment. Two studies (Durham 2003; Eckman 1992) attempted to keep patients from reporting their treatment groups to raters, however both reported that some patients disclosed allocation, and these were rated as unclear risk of bias. Stanton 1984 reported that some outcomes were assessed blindly and others were not. Hogarty 1997-study 1, Hogarty 1997-study 2, and Wirshing 1991 reported raters who were not blinded and were rated as high risk of bias.

Only Coyle 1988 and Stanton 1984 reported that they attempted to keep therapists blinded to the specific study hypothesis.

Incomplete outcome data

Overall, eighteen studies were rated as low risk of bias for incomplete outcome data. Lewis 2002b, Penn 2009, Sensky 2000b, Tarrier 1998, and Uzenoff 2007 reported outcomes with an intention-to-treat method, although Tarrier 1998 carried out an intention-to-treat analysis post-treatment only, and not at the one- or two-year follow-ups. Levine 1998 reported no participants leaving early. Sixteen studies reported reasons for losses to follow-up (Coyle 1988; Durham 2003; Falloon 1982; Haddock 1999; Hogarty 1997-study 1; Hogarty 1997-study 2; Kemp 1996; Klein 1998; Lewis 2002b; Malm 1982; O'Donnell 1999; Pinto 1999; Spaulding 1999; Tarrier 1998; Turkington 2000; Uzenoff 2007).

Of the five studies with unclear risk of bias for incomplete outcome data, Davidson 2004, Dincin 1982, Eckman 1992, Levine 1998, and Wirshing 1991 did not report on attrition. Only Stanton 1984 had a high risk of bias due to high rates of attrition reported without reasons.

Selective reporting

Five studies were of low risk of bias with regard to selective reporting (Davidson 2004, Lewis 2002b, Penn 2009, Sensky 2000b, and Uzenoff 2007). The remaining 19 studies were of high risk of bias, mainly due to poor data reporting - either they did not report data for outcomes or SDs were missing.

Other potential sources of bias

Four trials were unclear if they were subject to other biases (Coyle 1988; Haddock 1999; Levine 1998; Pinto 1999). All other trials appeared free from other sources of bias.



Effects of interventions

See: Summary of findings for the main comparison SUPPORTIVE THERAPY versus STANDARD CARE for schizophrenia; Summary of findings 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT for schizophrenia; Summary of findings 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY for schizophrenia; Summary of findings 4 SUPPORTIVE THERAPY versus FAMILY THERAPY for schizophrenia; Summary of findings 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION for schizophrenia

For dichotomous outcomes, we calculated risk ratios (RR) and for continuous outcomes, we calculated mean differences (MD), both with 95% confidence intervals (CI).

Comparison 1: SUPPORTIVE THERAPY versus STANDARD CARE

Five included studies compared supportive therapy with standard care (Coyle 1988; Davidson 2004; Durham 2003; Lewis 2002b; Tarrier 1998).

1.1 Global state: relapse

We found no significant difference in relapse rates between supportive therapy and standard care (Tarrier 1998, medium term, n = 54, RR 0.12 Cl 0.01 to 2.11; long term, n = 54, RR 0.96 Cl 0.44 to 2.11, Analysis 1.1). In this trial, relapses were defined as readmission to hospital for clinical deterioration that lased at least five days and resulted in functional impairment.

1.2 Service outcomes: hospitalisation

Hospitalisation rates did not differ significantly between people treated with supportive therapy and standard care (1 RCT, n = 48, RR 1.00 CI 0.07 to 15.08, Analysis 1.2).

1.3 Mental state: general

1.3.1 No clinically important improvement in general mental state

Medium-term data for 'no clinically important improvement in general mental state' were not statistically significant (Tarrier 1998, n = 54, RR 0.95 CI 0.77 to 1.17), and longer-term data (2 RCTs, n = 98, RR 0.95 CI 0.82 to 1.11) were also equivocal (Analysis 1.3).

1.3.2 Average endpoint general mental state score

Total scores on the PANSS were not significantly different at both short- (Lewis 2002b, n=131, MD -4.42 CI -10.1 to 1.29) and long-term time points (Durham 2003, n=36 MD 4.70 CI -6.71 to 16.11, Analysis 1.4).

1.4 Mental state: specific symptoms

1.4.1 Average endpoint negative symptoms score

Skewed data from the SANS scale are presented in tabular form (Analysis 1.5).

1.4.2 Average endpoint positive symptoms score

We found that the PANSS positive subscale data from the Lewis 2002b study showed no significant difference between supportive therapy and standard care (n = 131, MD -1.09 Cl -2.84 to 0.66, Analysis 1.6). One study measured severity of positive symptoms using selected items from the BPRS, but data are skewed and are presented as 'Other data' in a table (Analysis 1.7).

1.4.3 Average endpoint delusions score and hallucinations score

Skewed data from the PSYRATS scale 'delusions score' and 'hallucinations score' were obtained from one trial and are presented in a tabular form (Analysis 1.8; Analysis 1.9).

1.4.4 Average endpoint depressive symptoms score

Davidson 2004 found no statistical significance between the treatment groups for Center for Epidemiological Studies-Depression Scale (CESD) (n = 260, MD 1.61 CI -1.61 to 4.83, Analysis 1.10).

1.5 Leaving the study early

We found no significant difference in rates of attrition between supportive therapy and standard care (4 RCTs, n = 354, RR 0.86 CI 0.53 to 1.40, Analysis 1.11).

1.6 General functioning

1.6.1 Average endpoint in general functioning score

This was assessed by Durham 2003 using the GAS. This study found no significant difference between supportive therapy and standard care (n = 29, MD 1.40 CI -5.09 to 7.89, Analysis 1.13). Davidson 2004 used the GAF-M and also found no significant results (n = 260, MD -2.66 CI -6.20 to 0.88 Analysis 1.12).

1.6.2 Average endpoint in social functioning score

Davidson 2004 found no statistical significance between the treatment groups for social functioning measured on the SFS (n = 260, MD -0.67 CI -7.05 to 5.71, Analysis 1.14).

1.7 Satisfaction with treatment: recipient of care not satisfied with treatment

We found no significant difference in the numbers of recipients of care who were not satisfied with their treatment (Durham 2003, n = 44, RR 1.83 CI 0.75 to 4.47, Analysis 1.15).

1.8 Quality of life

1.8.1 Average endpoint self-esteem score

Davidson 2004 found no significant results between standard care and supportive therapy for self-esteem on the Rosenberg Self-Esteem Scale (RSES) (n = 260, MD -1.21 CI -2.85 to 0.43, Analysis 1.16).

1.8.2 Average endpoint well-being score

Davidson 2004 also assessed well-being on the WBS and found no significant difference in psychological functioning between treatment groups (n = 260, MD -2.73 CI -6.04 to 0.58, Analysis 1.17).

1.8.3 Average endpoint global health score

Global health was measured by Davidson 2004 on the GHQ and found no difference in nonpsychotic psychiatric symptomatology (n = 260, MD -2.45 CI -2.41 to 7.31, Analysis 1.18)

1.9 Death

The number of deaths occurring in the Tarrier 1998 study at medium term, were not significantly different (supportive therapy 1/26, standard care 0/28, n = 54, RR 3.22 Cl 0.14 to 75.75). Two trials reporting long-term data were also non-significant (n = 92, RR 2.87 Cl 0.31 to 26.63, Analysis 1.19).



1.10 Medication: no reduction in dose of antipsychotic

Durham 2003 reported on the number of people who had no reduction in the dose of antipsychotic medication at the endpoint of the trial. We found no statistically significant difference between the groups (n = 44, RR 0.81 CI 0.59 to 1.12, Analysis 1.20).

1.11 Costs

Most studies did not measure economic outcomes and no studies reported on costs in a way that could be added to the analysis. Klein 1998 reported that their findings suggest "that if the peer social supporter (the FC) had not been available, the study group may have needed as many inpatient days as the comparison group" and that this would have equated to "250 inpatient days (10 clients x 25 days)" that could have cost \$250,000 (based on average cost of \$1,000 per day, per client for inpatient psychiatric care) and compare this to "the cost of 10 FCs for 6 months, including administrative and programmatic costs, was about \$30,000".

Comparison 2: SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT

Twenty included studies compared supportive therapy against any other psychological or psychosocial treatment (Coyle 1988; Dincin 1982; Durham 2003; Eckman 1992; Falloon 1982; Haddock 1999; Hogarty 1997-study 1; Hogarty 1997-study 2; Kemp 1996; Levine 1998; Lewis 2002b; Penn 2009; Pinto 1999; Sensky 2000b; Spaulding 1999; Stanton 1984; Tarrier 1998; Turkington 2000; Uzenoff 2007; Wirshing 1991).

2.1 Global state: relapse

No significant difference in relapse rates were found in the supportive therapy and other therapies in the medium term (2 RCTs, n = 100, RR 2.86 CI 0.32 to 25.24, Analysis 2.1) or the long term (5 RCTs, n = 270, RR 1.19 CI 0.66 to 2.16, Analysis 2.1). Heterogeneity was high ($I^2 = 77\%$), so a random-effects model was used.

2.2 Global state: no remission

Falloon 1982 reported the number of people who did not experience remission of symptoms. We found a statistically significant advantage for other psychological therapies (in this case family therapy) over supportive therapy (n = 39, RR 1.87 CI 1.11 to 3.15, Analysis 2.2).

2.3 Service outcomes: hospitalisation

We found medium-term data for hospitalisation were equivocal (3 RCTs, n = 153, RR 1.60 CI 0.85 to 3.00), but longer-term data were statistically significant, favouring the other psychological or psychosocial therapies over supportive therapy (4 RCTs, n = 306, RR 1.82 CI 1.11 to 2.99, Analysis 2.3).

2.4 Mental state: general

2.4.1 No clinically important improvement

We found a significant difference favouring other therapies over supportive therapy for 'no clinically important improvement' over the long term (3 RCTs, n = 194, RR 1.27 Cl 1.04 to 1.54, Analysis 2.4), but not at medium-term follow-up (Tarrier 1998, n = 59, RR 1.27 Cl 0.95 to 1.70).

2.4.2 Average endpoint score

i) PANSS total

A significant difference in favour of other psychological treatment (in this case cognitive behavioural therapy (CBT)) was found for general mental state in the long term (2 RCTs, n = 105, MD 5.82 Cl 1.04 to 10.60, Analysis 2.5). However, in the short term, the data were highly heterogeneous ($l^2 = 98\%$). We have concerns about the data reported in Levine 1998, (see discussion section Quality of the evidence). We think the data in Levine 1998 may have been incorrectly reported or may have been an anomaly due to the small numbers in each treatment group. Also the standard deviations were very small, and we believe the authors may have erroneously reported standard errors as standard deviations. However, when we remove Levine 1998 from the analysis, heterogeneity is not restored and remains at 83%, therefore data were not pooled for this outcome.

ii) BPRS

No significant difference in mental state measured on the BPRS was found in the short term (2 RCTs, n=92, MD -1.07 CI -5.08 to 2.94); Pinto 1999 found a significant difference in favour of the control group (in this case CBT) in the long term (n=37, MD 7.60 CI 0.90 to 14.30, Analysis 2.6).

iii) BPRS short form

Kemp 1996 also used the short form of the BPRS, and found no significant differences at short- (n = 74, MD -0.90 CI -3.02 to 1.22), medium- (n = 67, MD 2.20 CI -1.18 to 5.58) or long-term follow-up (n = 45, MD 2.30 CI -0.54 to 5.14, Analysis 2.7).

iv) CPRS

One study presented skewed data from the CPRS (Sensky 2000b). We have reported these in a table (Analysis 2.8).

v) PANSS general subscale

No significant difference was found for general symptoms measured on the PANSS general subscale in the medium term (1 RCT, n=19, MD 2.86 CI -3.21 to 8.93, (Analysis 2.9); in the long term a significant difference was found in favour of the control group (1 RCT, n=65, MD 3.60 CI 0.84 to 6.36). For the short term, we did not pool the data as there was high heterogeneity ($I^2=97\%$) and we believe the data reported in Levine 1998 may have been incorrectly reported (see discussion section Quality of the evidence).

2.5 Mental state: specific symptoms

2.5.1 No clinically important improvement in negative and depressive symptoms

For the outcome of 'no clinically important improvement in negative symptoms', there was no significant difference between supportive therapy and other therapies (Sensky 2000b, n = 90, RR 0.95 CI 0.63 to 1.46 Analysis 2.10). Similarly, for the outcome of 'no important improvement in depressive symptoms', no significant differences were observed (Sensky 2000b, n = 90, RR 1.53 CI 0.92 to 2.55 Analysis 2.11). However, results show that people who underwent supportive therapy are more likely to have had an episode of affective symptoms than those who underwent other therapies (2 RCTs, n = 151, RR 1.84 CI 1.15 to 2.94, Analysis 2.12).



2.5.2 Average endpoint in negative symptoms

i) PANNS negative

No significant difference was found for general symptoms measured on the PANSS negative subscale in the medium term (1 RCT, n = 19, MD 1.70 CI -4.00 to 7.40) or the long term (1 RCT, n = 65, MD 0.30 CI -1.96 TO 2.56, Analysis 2.13). For the short term, we did not pool the data as there was high heterogeneity ($I^2 = 92\%$) and we believe the data reported in Levine 1998 may have been incorrectly reported (see discussion section Quality of the evidence).

ii) SANS

SANS medium-term data (Pinto 1999) were equivocal, Analysis 2.14. Another study also reported SANS data that are skewed, and are therefore reported separately in tabular form, Analysis 2.15.

2.5.3 Average endpoint in positive symptoms

i) PANSS positive

A significant difference in favour of other psychological treatment was found for positive symptoms in the long term (1 RCT, n = 65, MD 2.30 CI 0.60 to 0.40, Analysis 2.16). However, in the medium term no significant difference was found (1 RCT, n = 19, MD 2.71 CI -0.71 to 6.13). For the short term we did not pool data as they were highly heterogeneous ($I^2 = 83\%$). Furthermore, we have concerns about the data reported Levine 1998, (see discussion section Quality of the evidence). We think the data in Levine 1998 may have been incorrectly reported or may have been an anomaly due to the small numbers in each treatment group. Also the standard deviations were very small, and we believe the authors may have erroneously reported standard errors as standard deviations. However, when we remove Levine 1998 from the analysis, heterogeneity is not restored and remains at 78%.

ii) SAPS

One study reported data obtained from the SAPS. These data are skewed and are presented as 'Other data' in a table (Analysis 2.17).

iii) BPRS

One study reported data regarding severity of positive symptoms on the BPRS. However these data are skewed and are reported separately (Analysis 2.18).

2.5.4 Average endpoint in delusion, hallucination and voices symptoms

i) PANSS thought disturbance

Other therapies were found to significantly improve the PANSS thought disturbance cluster score (Levine 1998 n = 12, MD 4.30 Cl 1.17 to 7.43, Analysis 2.19, however, we believe these data may have been incorrectly reported (see above and discussion section Quality of the evidence).

ii) PSYRATS

Skewed data from the PSYRATS delusions score and hallucinations score are also presented in tabular form (Analysis 2.20; Analysis 2.21). Penn 2009 reported data from the voices score and found no significant difference at short term (n = 65, MD 0.10 CI -3.63 to 3.83) and long term (n = 65, MD 0.00 CI -4.76 to 4.76, Analysis 2.22).

iii) BAVQ

Penn 2009 reported skewed data on beliefs about voices using the BAVQ subscales, and are presented as 'Other data' in a table (Analysis 2.23).

2.5.5 Average endpoint in depressive symptoms

One study reported results from the MADRS and another from the BDI-II, but data are skewed and are reported in tabular form (Analysis 2.23 and Analysis 2.24, respectively).

2.6 Leaving the study early

We found no significant differences between supportive therapy and other treatments for leaving the study early (19 RCTs, n = 1412, RR 1.03 CI 0.87 to 1.21, Analysis 2.26). No difference was also found for general reasons for leaving the study early (17 RCTs, n = 1261, RR 0.97 CI 0.83 to 1.15). Treatment-related reasons for leaving the study early were significantly worse in the supportive therapy group (2 RCTs, n = 151, RR 2.15 CI 1.07 to 4.31).

2.7 General functioning

2.7.1 Average endpoint in general functioning

i) GAF & GAS

We found GAF scores (Kemp 1996) favoured CBT (short term, n = 70, MD -9.50 CI -16.11 to -2.89, and medium term, n = 67, MD -12.60 CI -19.43 to -5.77, Analysis 2.27). When we combined long term data from the GAF and GAS it was heterogeneous (I^2 = 84%) and so data were not pooled.

ii) SFS

Penn 2009 measured social functioning on the SFS, and found no significant difference in the short term (n = 65, MD -7.20 CI -17.86 to 3.46) and the long term (n = 65, MD -8.80 CI -21.67 to 4.07, Analysis 2.28).

2.7.2 Specific aspects

No advantage or disadvantages were found for people who had supportive therapy in terms of the number of people not in paid work (2 RCTs, n = 171, RR 1.03 CI 0.84 to 1.25, Analysis 2.29). As regards admission to residential placement (1 RCT, n = 39, RR 1.05 CI 0.24 to 4.59, Analysis 2.30) or admission to jail (1 RCT, n = 39, RR 1.05 CI 0.24 to 4.59, Analysis 2.31), similarly, we found no significant differences.

2.8 Satisfaction with treatment

One study looked at the proportion of people who were not satisfied with the care they received. We found a significant difference with participants in the other therapies group being less likely to be unsatisfied (Durham 2003, n = 45, RR 3.19 CI 1.01 to 10.7,, Analysis 2.32).

2.9 Quality of life

There was no significant difference in quality of life scores between the two groups (Uzenoff 2007, n = 19, MD -0.07 CI -21.11 to 20.97, Analysis 2.33).

2.10 Death

No significant differences were found for deaths during medium and long-term follow-up (Analysis 2.34).



2.11 Behaviour

A significant disadvantage was conferred on people who received supportive therapy in terms of social impairment measured on the SBAS (Falloon 1982, n = 39, RR 1.46 CI 1.04 to 2.04, Analysis 2.35). We found no significant effect on the number of people who showed a poor coping style with relatives (Falloon 1982, n = 39, RR 0.90 CI 0.37 to 2.20, Analysis 2.36).

2.12 Engagement with services

The rates of poor attendance at appointments were recorded in one study, and no significant differences were found between treatment groups (Falloon 1982, n = 39, RR 1.93 CI 0.89 to 4.17, Analysis 2.37.

2.13 Insight

Results from the SAI are skewed, and are presented in tabular form (Analysis 2.38).

2.14 Compliance

2.14.1 Poor compliance to therapy

We found compliance to be significantly worse amongst those who received supportive therapy than those in the control group (in this case, family therapy) (2 RCTs, n = 58, RR 2.63 CI 1.30 to 5.35, Analysis 2.39). However, by long-term follow-up, data were equivocal (1 RCT, n = 39, RR 1.29 CI 0.69 to 2.39).

2.14.2 Medication adherence (self-report)

No significant difference for self-reported medication adherence was found (Uzenoff 2007, n = 19, MD 1.00 CI 0.83 to 1.21, Analysis 2.40).

2.14.3 Average endpoint score (ROMI)

Uzenoff 2007 found no significant difference for medication adherence (n = 19, MD 0.45 CI -3.58 to 4.48, Analysis 2.41) or nonadherence (n = 19, MD 1.75 CI -0.61 to 4.11, Analysis 2.42).

2.15 Medication

Data from one small trial showed a significant difference, favouring other therapies for prescribed IM depot medication (Falloon 1982, n = 39, RR 2.11 CI 0.99 to 4.47, Analysis 2.43).

2.16 Attitude to medication

On the AMQ, the other therapies group scored significantly better than those allocated to supportive therapy (Kemp 1996, n = 74, MD -4.50 CI -6.83 to -2.17, Analysis 2.44). The DAI at short-term (Kemp 1996, n = 63, MD -5.70 CI -9.35 to -2.05, Analysis 2.45) and long-term follow-up (Kemp 1996, n = 44, MD -4.90 CI -9.38 to -0.42) also favoured the other therapies group.

2.17 Economic outcomes

The only study that investigated direct costs of the interventions produced skewed data, which are presented in a table (Analysis 2.46).

Comparison 3: SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY (CBT)

Thirteen included studies compared supportive therapy with CBT (Durham 2003; Haddock 1999; Hogarty 1997-study 1; Hogarty 1997-

study 2; Kemp 1996; Levine 1998; Lewis 2002b; Penn 2009; Pinto 1999; Sensky 2000b; Spaulding 1999; Tarrier 1998; Turkington 2000).

3.1 Global state: relapse

Relapse rates did not differ significantly between supportive therapy and CBT at medium term (2 RCTs, n = 100, RR 2.86 CI 0.32 to 25.24, Analysis 3.1). Longer-term data were heterogeneous ($I^2 = 78\%$) and so were not pooled.

3.2 Service outcomes: hospitalisation

Hospitalisations did not differ significantly between supportive therapy and CBT groups (medium term, 3 RCTs, n = 153, RR 1.60 CI 0.85 to 3.00, long term, 1 RCT, n = 65, RR 0.73 CI 0.18 to 3.00, Analysis 3.2).

3.3 Mental state: general

3.3.1 No clinically important improvement in general mental state

We found no significant difference for 'no clinically important improvement in general mental state' over the long term (3 RCTs, n=194, RR 1.23 CI 0.89 to 1.70), and at medium-term follow-up (1 RCT, n=59, RR 1.27 CI 0.95 to 1.70, Analysis 3.3).

3.3.2 Average endpoint general mental state score

i) PANSS total

A significant difference in general mental state between supportive therapy and CBT was demonstrated in the long term (2 RCTs, n = 105, MD 5.82 CI 1.04 to 10.60, Analysis 3.4). However, in the short term, the data were highly heterogeneous ($I^2 = 98\%$). We have concerns about the data reported Levine 1998, (see discussion section Quality of the evidence). We think the data in Levine 1998 may have been incorrectly reported or may have been an anomaly due to the small numbers in each treatment group. Also the standard deviations were very small, and we believe the authors may have erroneously reported standard errors as standard deviations. However, when we remove Levine 1998 from the analysis, heterogeneity is not restored and remains at 83% so data were not pooled for this outcome.

ii) BPRS

No significant difference in mental state measured on the BPRS was found in the short term (2 RCTs, n=92, MD -1.07 CI -5.08 to 2.94); Pinto 1999 found a significant difference in favour of CBT in the long term (n=37, MD 7.60 CI 0.90 to 14.30, Analysis 3.5).

iii) BPRS short form

Kemp 1996 also used the short form of the BPRS, and found no significant differences at short- (n = 74, MD -0.90 CI -3.02 to 1.22), medium- (n = 67, MD 2.20 CI -1.18 to 5.58) or long-term follow-up (n = 45, MD 2.30 CI -0.54 to 5.14, Analysis 3.6).

iv) CPRS

One study presented skewed data from the CPRS (Sensky 2000b). We have reported these in a table, Analysis 3.7.

v) PANSS general subscale

Levine 1998 reported PANSS general subscale data, which showed an advantage for other therapies over supportive therapy in the short term (n = 12, MD 17.10 CI 13.76 to 20.44), while Penn 2009



showed also showed a significant difference in favour of CBT in the short term (n = 65, MD 4.40 CI 1.36 to 7.44) and long term (n = 65, MD 3.60 CI 0.84 to 6.36, Analysis 3.8). We did not pool data for this outcome as the short-term results where highly heterogenous ($I^2 = 97\%$), and we believe the Levine 1998 data may have been incorrectly reported (see above and discussion section Quality of the evidence).

3.4 Mental state: specific symptoms

3.4.1 No clinically important improvement in negative and depressive symptoms

We found no significant difference in improvement in negative symptoms between supportive therapy and CBT (Sensky 2000b, n = 90, RR 0.95 CI 0.63 to 1.46, Analysis 3.9). Similarly, for depressive symptoms, we found no significant differences (Sensky 2000b, n = 90, RR 1.53 CI 0.92 to 2.55, Analysis 3.10). However, results show that people who underwent supportive therapy are more likely to have had an episode of affective symptoms than those who underwent CBT (2 RCTs, n = 101, RR 2.17 CI 1.16 to 4.06, Analysis 3.11).

3.4.2 Average endpoint in negative symptoms

i) PANSS negative

PANSS negative subscale data reported by Levine 1998 significantly favoured CBT in the short term (n = 12, MD 7.20 CI 3.78 to 10.62), while Penn 2009 showed no significance in the short term (n = 65, MD 0.30 CI -1.55 to 2.15) and long term (n = 65, MD 0.30 CI -1.96 to 2.56, Analysis 3.12). The short-term data were highly heterogenous ($I^2 = 92\%$) and we have concerns about Levine's data (see above and discussion section Quality of the evidence) and so data were not pooled.

ii) SANS

Pinto 1999 reported SANS data and found no difference between treatment groups (n = 37, MD 6.60 CI -5.81 to 19.01). Other skewed SANS data for Sensky 2000a and Tarrier 1998 are reported separately in tabular form Analysis 3.14.

3.4.3 Average endpoint in positive symptoms

i) PANSS positive

Three studies evaluated positive symptoms using the PANSS positive subscale, and no significant difference was found Lewis 2002b and a significant difference in favour of CBT was found in Levine 1998 and Penn 2009. The pooled data were highly heterogenous ($I^2 = 83\%$); when we removed Levine 1998, for which we have concerns about the data reported (see above and discussion section Quality of the evidence), the heterogeneity remain high with $I^2 = 78\%$ and so data were not pooled (Analysis 3.15). In the long term, Penn 2009 favoured CBT (n = 65, MD 2.30 CI 0.60 to 4.00, Analysis 3.15).

ii) SAPS

Tarrier 1998 reported data obtained from the SAPS, but data are skewed and presented as 'Other data' in a table (Analysis 3.16).

iii) BPRS

Skewed positive symptoms data from the BPRS were obtained from Tarrier 1998, and are presented as 'Other data' in a table (Analysis 3.17).

3.4.4 Average endpoint in delusion, hallucination and voices symptoms

i) PANSS thought disturbance

Cognitive behavioural therapy did not improve scores on the PANSS thought disturbance cluster significantly more than supportive therapy (Levine 1998 n = 12, MD 4.30 Cl 1.17 to 7.43, Analysis 3.18), however, we believe these data may have been incorrectly reported (see above and discussion section Quality of the evidence).

ii) PSYRATS

Skewed data from the PSYRATS scale, delusions score and hallucinations score are also presented in tabular form (Analysis 3.19; Analysis 3.20). Penn 2009 reported data from the voices score and found no significant difference at short term (n = 65, MD 0.10 CI -3.63 to 3.83) and long term (n = 65, MD 0.00 CI -4.76 to 4.76, Analysis 3.21).

iii) BAVQ

Penn 2009 reported skewed data on beliefs about voices using the BAVQ subscales, and are presented as 'Other data' in a table (Analysis 3.22).

3.4.5 Average endpoint in depressive symptoms

Sensky 2000b reported results from the MADRS and Penn 2009 from the BDI-II, but data are skewed and are reported in tabular form (Analysis 3.23 and Analysis 3.24, respectively).

3.5 Leaving the study early

We found no significant differences (general reasons) for leaving the study early, between supportive therapy (10 RCTs, n = 711, RR 0.93 CI 0.66 to 1.30, Analysis 3.25) and those allocated to CBT. For those who left due to treatment-related reasons data were again non-significant (2 RCTs, n = 101, RR 2.34 CI 0.90 to 6.10), but heterogeneous ($l^2 = 66\%$).

3.6 General functioning

We found GAF scores (Kemp 1996) favoured CBT (short term, n = 70, MD -9.50 CI -16.11 to -2.89, and medium term, n = 67, MD -12.60 CI -19.43 to -5.77, Analysis 3.26). When we combined long-term data from the GAF and GAS they were heterogeneous ($I^2 = 84\%$) and so data were not pooled.

Penn 2009 measured social functioning on the SFS, and found no significant difference in the short term (n = 65, MD -7.20 CI -17.86 to 3.46) and the long term (n = 65, MD -8.80 CI -21.67 to 4.07, Analysis 3.27).

3.7 Satisfaction with treatment

Durham 2003 looked at the proportion of people who were not satisfied with the care they received. We found a significant difference between treatment groups, with people receiving CBT less likely to be unsatisfied (n = 45, RR 3.19 CI 1.01 to 10.07, Analysis 3.28).

3.8 Quality of life

One study found no significant results for the RSES (Penn 2009, n = 65 short term MD -0.80 CI -3.77 to 2.17; long term MD -1.70 CI -5.19 to 1.79, Analysis 3.29).



3.9 Death

Two deaths occurred in Tarrier 1998, one from each intervention group, and one death occurred in the Durham 2003 study from the supportive therapy group (Analysis 3.30).

3.10 Insight

Results from the Schedule for Assessment of Insight are skewed, and are presented in tabular form Analysis 3.31, as are results from the Beck Cognitive Insight Scale, Analysis 3.32.

3.11 Medication

There was no significant difference for no reduction in doses of antipsychotic medication between the CBT and supportive therapy groups (Sensky 2000b, n = 90, MD 0.89 CI 0.68 to 1.17, Analysis 3.33).

3.12 Attitude to medication

On the AMQ, people who were allocated to CBT scored significantly better than those allocated to supportive therapy (Kemp 1996, n = 74, MD -4.50 CI -6.83 to -2.17, Analysis 3.34). Similarly, the DAI at short-term (Kemp 1996, n = 63, MD -5.70 CI -9.35 to -2.05, Analysis 3.35) and long-term follow-up (Kemp 1996, n = 44, MD -4.90 CI -9.38 to -0.42) also significantly favoured CBT compared with supportive therapy.

3.13 Economic outcomes

The only study investigating direct costs of the interventions produced skewed data and is reported in tables (Analysis 3.36).

Comparison 4: SUPPORTIVE THERAPY versus FAMILY THERAPY

Two included studies compared supportive therapy with family therapy (Falloon 1982; Hogarty 1997-study 1).

4.1 Global state: relapse and remission

Two trials reported relapse rates, but these showed heterogeneity ($I^2 = 87.1\%$) and so data were not pooled (Analysis 4.1). We found participants in the Falloon 1982 study were more likely to experience remission of their symptoms if they had family therapy rather than supportive therapy (n = 39, RR 1.87 CI 1.11 to 3.15, Analysis 4.2).

4.2 Service outcome: hospitalisation

No significant differences were found in hospitalisation rates between the treatment groups (Falloon 1982, n = 39, RR 1.93 CI 0.89 to 4.17, Analysis 4.3).

4.3 Mental state: episode of affective symptoms

Data were only available for one outcome relating to mental state - the likelihood of having an episode of affective symptoms. We found no significant difference between family therapy and supportive therapy (Hogarty 1997-study 1, n = 48, RR 1.71 CI 0.82 to 3.60, Analysis 4.4).

4.4 Leaving the study early

We found no significant difference for the number of participants leaving the study early for general reasons (Falloon 1982, n = 39, RR 0.70 CI 0.13 to 3.75) and treatment-related reasons (Hogarty 1997-study 1, n = 48 RR 1.60 CI 0.61 to 4.19, Analysis 4.5).

4.5 General functioning

People who had supportive therapy were no more or less likely to have failed to find paid work than people who had family therapy (Falloon 1982, n = 39, RR 0.96 CI 0.57 to 1.63, Analysis 4.6). Rates of admission to residential placement (Falloon 1982, n = 39, RR 1.05 CI 0.24 to 4.59, Analysis 4.7) and admission to jail (1 RCT, n = 39, RR 1.05 CI 0.24 to 4.59, Analysis 4.8) did not differ significantly between treatment groups.

4.6 Behaviour

Falloon 1982 assessed the frequency of social impairment using the Social Behaviour Adjustment Schedule. We found social impairment to be significantly more frequent in the supportive therapy intervention compared with family therapy (n = 39, RR 1.46 CI 1.04 to 2.04, Analysis 4.9). The number of people displaying a poor coping style with relatives did not differ significantly between groups (Falloon 1982, n = 39, RR 0.90 CI 0.37 to 2.20, Analysis 4.10).

4.7 Engagement with services

The proportion of people who were poor attendees at appointments was not significantly different (Falloon 1982, n = 39, RR 1.93 CI 0.89 to 4.17, Analysis 4.11) between groups.

4.8 Compliance

We found compliance to be significantly worse amongst those who received supportive therapy than those in the family therapy group (Falloon 1982, n = 39, RR 2.63 CI 1.30 to 5.35, Analysis 4.12). However, by long term follow-up data were equivocal (n = 39, RR 1.29 CI 0.69 to 2.39).

4.9 Medication

The difference in frequency of prescription of antipsychotic depot medication between groups was not statistically significant (n = 39, RR 2.11 CI 0.99 to 4.47, Analysis 4.13).

Comparison 5: SUPPORTIVE THERAPY versus PSYCHOEDUCATION

Two included studies compared supportive therapy with psychoeducation (Coyle 1988; Uzenoff 2007).

5.1 Service outcomes: hospitalisation

No significant differences in hospitalisation rates were found (Coyle 1988, n = 47, RR 0.48 CI 0.05 to 4.93, Analysis 5.1).

5.2 Mental state

5.2.1 No clinically important improvement in general mental state

Uzenoff 2007 found no significant difference between supportive therapy and psychotherapy for 'no clinically important improvement in general mental state' (n = 19, MD 1.61 CI 0.96 to 2.68, Analysis 5.2).

5.2.2 Average endpoint general symptoms score

There was no significant difference for PANSS general scores between the groups (n = 19, MD 2.86 CI -3.21 to 8.93, Analysis 5.3).



5.2.3 Average endpoint negative symptoms score

PANSS negative subscale data from Uzenoff 2007 showed no significant difference between supportive therapy and psychoeducation (n = 19, MD 1.70 Cl -4.00 to 7.40, Analysis 5.4).

5.2.4 Average endpoint positive symptoms score

PANSS positive subscale data from Uzenoff 2007 study showed no significant difference between supportive therapy and psychotherapy (n = 19, MD 2.71 CI -0.71 to 6.13, Analysis 5.5).

5.2.5 Average endpoint depression score

No significant difference was found for CDRS between supportive therapy and psychotherapy (Uzenoff 2007, n = 19, MD 1.47 CI -1.35 to 4.29, Analysis 5.6)

5.3 Leaving the study early

We found no significant difference in the number of people leaving the study early between treatment groups (2 RCTs, n = 71, RR 0.57 CI 0.21 to 1.54, Analysis 5.7).

5.4 Quality of Life

There was no significant difference in quality of life scores between the two groups (Uzenoff 2007, n = 19, MD -0.07 CI -21.11 to 20.97, Analysis 5.8).

5.5 Death

Death rates were not significantly affected by treatment received (Coyle 1988, n = 47, RR 2.88 CI 0.12 to 67.29, Analysis 5.9).

5.6 Behaviour

Uzenoff 2007 found no significant differences in the composite scores from the ROMI and ITAQ for need for treatment (n = 19, MD -0.02 CI -0.44 to 0.40) and the benefits of medication (n = 19, MD -0.19 CI -0.56 to 0.18, Analysis 5.10).

5.7 Insight

No significant difference in insight into treatment scores were found between supportive therapy and psychotherapy (Uzenoff 2007, n = 19, MD -1.55 CI -5.85 to 2.75, Analysis 5.11)

5.8. Compliance

5.8.1 Poor compliance to therapy

All participants in Uzenoff 2007 attended at least six of the 14 sessions of therapy (Analysis 5.12).

5.8.2 Medication adherence (self-report)

No significant difference for self-reported medication adherence was found (Uzenoff 2007, n = 19, MD 1.00 CI -0.83 to 1.21, Analysis 5.13).

5.8.3 Average endpoint score (ROMI)

Uzenoff 2007 found no significant difference for medication adherence (n = 19, MD .45 CI -3.58 to 4.48, Analysis 5.14) or nonadherence (n = 19, MD 1.75 CI -0.61 to 4.11, Analysis 5.15) between supportive therapy and psychotherapy.

Comparison 6: SUPPORTIVE THERAPY versus REHABILITATION PROGRAMME

One study looked at supportive therapy compared to a rehabilitation programme (Dincin 1982).

6.1 Service outcomes

Participants in the rehabilitation group were significantly less likely to be hospitalised than those in the supportive therapy group (Dincin 1982, n = 132, RR 2.71 CI 1.22 to 6.02, Analysis 6.1).

6.2 Leaving the study early

There were no significant differences in the numbers who left the study early (Dincin 1982, n = 132, RR 1.45 CI 0.92 to 2.29, Analysis 6.2).

6.3 General functioning: no gainful employment

We found that the therapeutic intervention did not have a significant impact on the number of people not in gainful employment (Dincin 1982, n = 132, RR 1.04 CI 0.85 to 1.29, Analysis 6.3).

Comparison 7: SUPPORTIVE THERAPY versus SKILLS TRAINING

Three included studies compared supportive therapy with skills training (Coyle 1988; Eckman 1992; Wirshing 1991).

7.1 Service outcomes: hospitalisation

Hospitalisation rates were not found to differ significantly between supportive therapy and skills training (Coyle 1988, n = 47, RR 0.96 CI 0.06 to 14.43, Analysis 7.1).

7.2 Leaving the study early

Attritions rates showed no significant difference between treatment groups (3 RCTs, n = 168, RR 1.01 Cl 0.61 to 1.67, Analysis 7.2).

7.3 Death

Coyle 1988 reports data on death rates, with no significant difference between groups (n = 47, RR 2.88 CI 0.12 to 67.29, Analysis

Comparison 8: SUPPORTIVE THERAPY versus PSYCHODYNAMIC PSYCHOTHERAPY

Stanton 1984 compared supportive therapy with psychodynamic psychotherapy.

8.1 Leaving the study early

In the medium term, people in the psychodynamic therapy group were more likely to leave the study early than those in the supportive therapy arm (1 RCT, n = 164, RR 0.62 CI 0.42 to 0.91). However, by long term follow-up, the differences were no longer significant (Stanton 1984, n = 164, RR 0.89 CI 0.73 to 1.09, Analysis 8.1).

Comparison 9: SUPPORTIVE THERAPY versus COMBINATION OF OTHER PSYCHOSOCIAL INTERVENTIONS

Hogarty 1997-study 1 compared supportive therapy against a combination of other psychosocial interventions.



9.1 Global state: relapse

We found no significant difference for relapse rates between supportive therapy and those receiving a combination of other therapies, in this case CBT and family therapy (Hogarty 1997-study 1, n = 50, RR 1.48 CI 0.86 to 2.55, Analysis 9.1).

9.2 Mental state

The risk of having an episode of affective symptoms did not differ significantly between groups (Hogarty 1997-study 1, n = 50 RR 1.63 CI 0.81 to 3.28, Analysis 9.2).

9.3 Leaving the study early

The number of people who left the study early for treatment-related reasons were significantly higher in the supportive therapy group than in the combination group (Hogarty 1997-study 1, n = 50, RR $8.67\ Cl\ 1.17\ to\ 64.26$, Analysis 9.3).

Comparison 10: SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus CLIENT-FOCUSED CASE MANAGEMENT

Klein 1998 and O'Donnell 1999 compared supportive therapy and client-focused case management against client-focused case management alone.

10.1 Global state: relapse

The number of participants experiencing relapse did not differ between groups (Klein 1998, n = 61, RR 0.32 CI 0.05 to 2.14 Analysis 10.1).

10.2 Leaving the study early

The frequency of participants leaving the study early was not significantly different (2 RCTs, n = 145, RR 2.38 CI 1.15 to 4..3,9 Analysis 10.2).

10.3 Death

Death rates did not differ significantly between the two treatment groups (O'Donnell 1999, n = 84, RR 2.61 CI 0.11 to 62.26, Analysis 10.3).

Comparison 11: SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT Versus STANDARD CASE MANAGEMENT

O'Donnell 1999 compared supportive therapy and client-focused case management against standard case management.

11.1 Leaving the study early

There were no significant differences between the treatment and control group (O'Donnell 1999, n = 80, RR 0.88 CI 0.52 to 1.51, Analysis 11.1).

11.2 Death

We found no significant differences in death rates (O'Donnell 1999, $n=80, RR\ 2.35\ CI\ 0.10$ to 55.94 Analysis 11.2).

Comparison 12: SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING

Malm 1982 compared supportive therapy and skills training with skills training alone.

12.1 Global state

Relapse rates were not significantly different between the two treatment regimes (Malm 1982, n=80, RR 1.00 CI 0.49 to 2.04, Analysis 12.1). Similarly, we found no significant difference in the number of people who did not experience remission during the follow-up period (Malm 1982, n=80, RR 0.78 CI 0.54 to 1.12, Analysis 12.2).

12.2 Service outcomes

Data were available for the number of people who were not discharged from hospital during the follow-up period (all participants were inpatients at the start of the study). No significant differences were found between the two treatment groups (Malm 1982, n = 80, RR 1.14 Cl 0.46 to 2.85, Analysis 12.3).

12.3 Mental state

Data were only reported for two out of 45 items on the CPSRS, 'inability to feel' and 'derealisation'. For 'inability to feel', results showed no significant difference between the two treatment groups (Malm 1982, n = 80, MD 0.10 CI -0.08 to 0.28, Analysis 12.4). Data for 'derealisation' are skewed, and are presented as 'Other data' in a table (Analysis 12.5).

12.4 Leaving the study early

No significant difference was found for this outcome (Malm 1982, n = 80, RR 1.00 CI 0.35 to 2.84, Analysis 12.6).

12.5 General functioning

Data from the KAS were reported for the subscale entitled 'free-time activities' and for the syndrome entitled 'withdrawal'. For 'free-time activities', we found a statistically significant advantage for supportive therapy plus skills training, compared with skills training alone (Malm 1982, n = 80, MD 0.10 CI 0.02 to 0.18, Analysis 12.8). Data for 'withdrawal' are skewed, and are presented as 'Other data' in a table (Analysis 12.8).

12.6 Death

No significant differences were found in death rates between the two treatment groups (Malm 1982, n = 80, RR 2.00 CI 0.19 to 21.18, Analysis 12.9). Three deaths occurred, all of which were due to suicide.

DISCUSSION

Summary of main results

1. Supportive therapy versus standard care

There was a lack of data for this comparison, with only five studies included. Overall, the quality of the evidence was very low for the outcomes in the Summary of findings for the main comparison.

We found no evidence of significant differences between supportive therapy and standard care for rates of relapse and hospitalisation, clinical improvement in mental state, leaving the study early and quality of life and no studies reported data on general functioning and satisfaction with treatment.



2. Supportive therapy versus any other psychological or psychosocial treatment

Twenty studies were included in the comparison, which combines data from the comparisons for other psychological interventions, including cognitive behavioural therapy (CBT), family therapy and the quality of the evidence was very low for all outcomes except leaving the study early, which we rated as moderate in the Summary of findings 2.

For the outcomes rate of hospitalisations, clinical improvement in mental state and satisfaction of treatment for the recipient of care, we found evidence favouring any other psychological or psychosocial treatment. We found no evidence of significant differences for rate of relapse, leaving the study early and quality of life

3. Supportive therapy versus cognitive behavioural therapy (CBT)

Thirteen studies compared supportive therapy with CBT. Again, the quality of the evidence was very low for all outcomes except leaving the study early, which we rated as moderate (see Summary of findings 3).

We found no evidence of significant differences between supportive therapy and CBT for rates of relapse and hospitalisation, clinical improvement in mental state, leaving the study early and quality of life. For general functioning, we could not pool data as it showed high heterogeneity; one study showed equivocal data, the other found in favour of supportive therapy.

4. Supportive therapy versus family therapy

There was a lack of data for this comparison, with only two studies included. Overall, the quality of the evidence was very low for the outcomes in the Summary of findings 4.

We were not able to pool the data for rate of relapse due to high heterogeneity; one study found no difference in relapse rates, the other study found a difference in favour of family therapy. We found no differences for the rate of hospitalisation and leaving the study early. As there were limited data for this comparison, we had to use proxy data for two outcomes: for mental state we had data for participants experiencing an episode of affective symptoms and for general functioning we had data for participants with no paid work. Neither of the studies reported data for satisfaction with treatment and quality of life.

5. Supportive therapy versus psychoeducation

Again, we were only able to include two studies in this comparison and the quality of the evidence was very low (see Summary of findings 5).

We found that there was no evidence of significant differences between supportive therapy and psychoeducation for the outcomes with data – rate of hospitalisation, no clinical improvement in mental state and leaving the study early. No studies reported data for rate of relapse, general functioning, satisfaction with treatment and quality of life.

Overall completeness and applicability of evidence

1. Few data

It is important to point out that the findings in this review are significantly weakened by the limited amount of available data. The search strategy identified five trials, involving 822 people, comparing supportive therapy with standard care. However, few data regarding the predetermined primary outcomes of interest could be extracted. The important clinical question, whether supportive therapy confers any advantage over standard care in the treatment of people with schizophrenia, remains unanswered. It is important to answer this question before results of research comparing supportive therapy with other psychological or psychosocial therapies can be interpreted fully and meaningfully.

More studies compared supportive therapy with other psychological or psychosocial interventions. There were, however, many comparison interventions. The number of studies comparing supportive therapy with each intervention was generally small, the main exception being CBT, for which we identified 13 studies.

2. Delivery of therapy

In trials of psychological interventions, there is question about whether the same therapists should provide two different interventions, potentially allowing factors such as level of experience of the therapists and individual differences in personality to be evenly distributed between groups. However, an alternative, and in our opinion more persuasive, argument is that different therapists should provide different therapies. This takes into account the likelihood that therapists have a loyalty to, and training and experience in, one particular type of therapy. This may be particularly important if they have been involved in generating the hypotheses which are being tested. The majority of studies in this review used the same therapists for supportive therapy and other psychological interventions. Most studies did not specify what training the therapists had received or what level of experience they had. Where studies did report details of therapist training, therapists who delivered supportive therapy were sometimes trained in other modalities, such as CBT, but not in supportive therapy. The studies which described standardised supportive therapy, with the use of a manual, were in the minority; as were the studies that evaluated or monitored adherence to the treatment model.

Not all studies attempted to match the amount of therapist contact. For example, Coyle 1988 compared supportive therapy with social skills training and psychoeducation. Supportive therapy sessions, however, were half as long as other therapy sessions, and were delivered individually rather than in a group.

More recent studies, such as Lewis 2002b, Penn 2009 and Sensky 2000b are of a much better quality. It seems likely that methodology and reporting have improved considerably over recent years, which will be valuable for future versions of this review. The concerns described above mean that only tentative conclusions can be drawn from this review.

3. Applicability

All trials used entry criteria to define who would be included in the study. These, however, varied from study to study. The studies involved people with schizophrenia and schizophrenia-like



illnesses defined by a range of criteria. Most studies, but not all, used some form of diagnostic criteria. Settings also varied, and both inpatients and outpatients were included. This suggests that the results would be valid for people with a diagnosis of schizophrenia or a related psychotic illness, whether they are being treated as an inpatient or on an outpatient basis. However, the possibility that patients typically studied in randomised controlled trials may differ in important ways from patients typically seen in mental health services should also be considered (Roth 1996). The results of this review should ideally be considered alongside results of effectiveness research. In effectiveness research, generalisability may be higher that for randomised controlled trials, although internal validity is lower (Margison 2000).

One aspect of trial setting which might be particularly important is the country in which trials are conducted. In this review, the majority were set in the USA. Seven were set in the UK, and one each in Israel, Italy, Sweden and Australia. There may be a difference in what is regarded as supportive therapy in the USA and in the UK (Holmes 1995), with, for example, supportive therapy in the USA perhaps being more frequent and more closely related to a psychodynamic approach than in the UK.

One of the difficulties in attempting this systematic review has been finding a clinically useful definition of supportive therapy. Despite the lack of a universally accepted definition, we felt it was important to define the intervention and identify the available evidence. Throughout the process, we have been aware of the risk that interventions included in the review would be too diverse. Readers of this review need to know what is meant by supportive therapy, and to be able to apply the findings in clinical practice. The included interventions do vary, and we have some concerns about whether they have enough in common to be grouped together in a useful way. The wide variation in the duration and frequency of sessions could be taken as evidence of differences between interventions in the included studies. Nevertheless, we note the lack of research in this field, and the lack of specific meta-analyses on supportive therapy. We feel the use of a broad definition of supportive therapy in this review is justified as producing an initial analysis of the available data. It is possible that narrower definitions of supportive therapy, perhaps used in more specific settings, could produce different results. For example, it may be possible that an untrained support worker within a community team could reduce the input required for a patient from other members of the team, or may increase the leisure and community activities of a patient. In this review we are unable to make comments on specific questions such as these.

Treatment manuals, training of therapists and monitoring of treatment fidelity, all help to enhance internal validity of trials. These were used in a minority of trials in this review. However, it can be argued that these measures may increase the apparent effectiveness of interventions over what is possible to reproduce in clinical practice. In many reports, little description of the supportive intervention was provided.

Two studies in particular described interventions which are different from the other supportive interventions. Kemp 1996 reports that therapists delivering supportive therapy were specifically instructed to decline to discuss treatment with people. O'Donnell 1999 also describes a somewhat different intervention. In this study, supportive therapy is more like a client-focused case management with advocacy. Advocacy, which comes within

our definition of supportive therapy, was provided by consumer advocates, who could be people who had experienced mental illness, relatives or carers of people with mental illnesses, or interested lay people. No particular frequency or duration of sessions was specified.

Quality of the evidence

There was little consensus between studies over which outcome variables to use. Definition of outcomes such as relapse also varied between trials.

Sample sizes were small for most studies, perhaps resulting in true beneficial or harmful effects going undetected.

The quality of the available data is also a cause for concern which needs to be borne in mind when interpreting the results of this review. Twenty of the 24 studies did not adequately report the use of allocation concealment, raising concerns about selection bias. We suspect that this would be most likely to act in the direction of biasing results against supportive therapy, as supportive therapy is used as a control treatment in the majority of studies in this review. Although double-blind trials are not possible when testing a psychological intervention, it is possible for therapists to be blinded to the study hypothesis. Only two studies attempted this. Not all studies rated outcomes blindly, resulting in a risk of detection biases. Attrition bias cannot be discounted either, as losses to follow-up were generally poorly reported, and intention-to-treat analyses were the exception rather than the rule.

These potential biases are particularly important in this review. This is because studies are almost all designed with supportive therapy as a comparator for other treatments, rather than being the primary treatment of interest to investigators. The hypothesis and the expectation of investigators is therefore likely to be that supportive therapy will not perform as well as the other treatment under investigation.

Heterogeneity was present, as judged by visual inspection and the I² statistic, for the outcomes: mental state measured on the Positive and Negative Syndrome Scale (PANSS), general functioning on the Global Assessment of Function disability scale (GAF) and Global Assessment Scale (GAS) when supportive therapy was compared with other psychological interventions or CBT alone; relapse when supportive therapy was compared with other psychological interventions CBT alone and family therapy; and leaving the study early when supportive therapy was compared with CBT. This may be due in part to the diversity of the interventions. There is a possibility that our attempt to investigate the effects of supportive therapy may not have been as successful as we might have hoped due to the variety of interventions meeting our criteria for supportive therapy or care. However, as high rates of heterogeneity were not found for other outcomes, this does not seem likely to be a major problem. Another possibility, and probably one that is more likely, is that the heterogeneity was due to diversity of control interventions (family therapy or CBT). Although homogeneity was not restored when data from Levine 1998 were excluded from the pooled analyses (due to the presence of heterogeneity). We think the data in Levine 1998 may have been incorrectly reported or may have been an anomaly due to the small numbers in each treatment group (n = 7). Also the standard deviations were very small, and we believe the authors may have erroneously reported standard errors as standard deviations.



Potential biases in the review process

1. The limitations of randomised trials of psychotherapy

In doing this review, we explicitly aimed to include only randomised controlled trials. However, it is important to recognise that evidence from randomised controlled trials is not the only form of evidence which can be used to guide clinical practice. There are particular problems with randomised controlled trials for psychological therapies. Psychotherapy relies on the uniqueness of the clinician-patient relationship, and ways of measuring outcomes which take account of this need to be developed (Holmes 2000). Randomised controlled trials employ techniques to enhance internal validity, such as frequent monitoring, the use of therapy manuals and excluding patients with dual diagnoses. These techniques may increase the apparent effectiveness of the intervention above what would routinely be achieved in clinical practice (Roth 1996). In randomised controlled trials of long-term therapy, there is a risk of contamination by other treatments (Margison 2000).

2. Definition of supportive therapy

Distinguishing supportive therapy from standard treatment has also been an issue in this review. We have largely relied on authors' descriptions of the interventions, and have classified some interventions described as treatment as usual as supportive therapy. Similarly, some interventions which were described as supportive in nature were not felt to meet the pre-determined inclusion criteria for this review. Descriptions have not always been of sufficient detail to judge whether a control intervention should be classified as supportive therapy or not, leaving the possibility that some data were in appropriately omitted. The lack of differences in outcomes between supportive therapy and standard care could suggest that this distinction is not a valid one. However, this is not the only interpretation of the results, and a real difference in effectiveness could exist, but may not have been identified due to paucity of data.

3. Strengths

Despite the reservations and difficulties outlined above, it is important that this review has been attempted. Supportive therapy is commonly used in clinical practice for people with schizophrenia. It may be cheaper than other therapies, such as CBT but is probably more costly and intensive than what is generally provided as standard care in many places. We also feel it to be a prevalent approach so it is important to systematically examine the effects of supportive therapy as a specific intervention. As far as we are aware, the only published study previously to have attempted this was the meta-analysis undertaken by the Department of Health for the purposes of the NICE Guideline on Schizophrenia (NICE 2003). The scope of this systematic review is not quite as focused as this work and therefore the NICE 2003 review is larger with more studies included. We feel our tighter definitions of supportive therapy are justified, workable and give a clearer picture of the evidence.

Agreements and disagreements with other studies or reviews

We know of no other reviews on supportive therapy.

AUTHORS' CONCLUSIONS

Implications for practice

1. For people with schizophrenia

The evidence presented in this review does not allow us to conclude that supportive therapy has any substantial benefit over standard care. Although there is no evidence of a treatment effect when comparing supportive therapy with standard care, the small number and small size of trials means that a treatment effect, either favouring supportive therapy or standard care, cannot be ruled out. This rather unsatisfactory conclusion also applies to other comparisons where supportive therapy is directly compared with several other psychological therapies, including cognitive behavioural therapy (CBT). At present there is not really conclusive evidence for people with schizophrenia to choose one of the other therapies over a more simple supportive approach.

2. For clinicians

Guidance encourages use of family and cognitive therapy for people with schizophrenia. From this review, one cannot conclude that supportive therapy offers anything different to other forms of psychological intervention. For some outcomes, including hospitalisation, general mental state and affective symptoms, data suggest an advantage for other therapies over supportive therapy but due to the small amount of data and the lack of consistency between outcomes measuring similar variables, firm conclusions should not be drawn from these results.

3. For mangers and policy makers

Unfortunately, the paucity of data means no clear recommendations for clinical practice can be made on the basis of this review. At present, there is no evidence from this review of a beneficial effect for supportive therapy over standard care, although such a beneficial or harmful effect cannot be ruled out. There is some evidence to suggest a disadvantage of supportive therapy in comparison with other therapies but this is scarce and is often derived from studies that may have some inherent bias towards the other therapies. Certainly, these data require replication in large, clinically meaningful randomised controlled trials before they can be used to guide clinical practice. It is important to highlight the lack of data regarding harmful or adverse effects of supportive therapy. This would be deemed unacceptable in a trial relating to a specific medication.

Implications for research

1. General

As with similar studies, public registration of a study before anyone is randomised would ensure that participants could be confident that people would know that the study had at least taken place. Unique study numbers would help researchers to identify single studies from multiple publications and reduce the risk of duplicating the reporting of data. Compliance with CONSORT (Moher 2001), both on the part of authors and editors, would help to clarify methodology and many outcomes. Failure to comply with such standard guidelines results in both loss of data and confusion in the results, neither of which help clinicians, patients or managers.



Intention-to-treat analysis should be performed on all outcomes and all trial data should be made easily accessible. A minimal requirement should be that all data should, at least, be presented as numbers. In addition, continuous data should be presented with means, standard deviations (or standard errors) and the number of participants. Data from graphs, 'P' values of differences and statements of significant or non-significant differences are of limited value. Unfortunately, in spite of the large numbers of participants randomised, we were unable to use most of the data in the trials included in this review due to the poor data reporting.

2. Specific

2.1 Reviews

Many of the excluded studies suggest new comparisons relevant to future reviews (Characteristics of excluded studies), too many for this exhausted group of systematic review authors to currently list.

2.2 Trials

Supportive therapy can be usefully investigated using randomised controlled trials. Of course, evidence from randomised controlled trials should be considered alongside evidence from other forms of research, but larger trials do seem warranted and preferably undertaken by those with clear equipoise. A suggestion for the design is outlined in Table 1.

2.2.1 Methods

Allocation concealment is vital in the design of future randomised controlled trials to minimise bias. The randomisation process should also be described fully. It is probably not possible to use double-blind methodology in trials of psychological and psychosocial interventions. However, bias can be minimised by ensuring therapists and people are blind to the specific study hypothesis where feasible, and by using blinded or independent outcome raters.

2.2.2 Participants

We would suggest that inclusion criteria for participants in such a study would be broad and that the therapy be given in the context of everyday practice.

2.2.3 Interventions

2.2.3.1 Supportive therapy

Future trials should clearly explain whether practitioners who deliver supportive therapy have been specifically trained, and if so

how. It may make the results more applicable if the therapists are trained but in the context of routine career development, rather than specific highly-trained specialised practitioners. In addition, using a therapy manual or protocol and attempting to ensure adherence to the model may at least increase internal validity of the trial. Ideally, different therapists should be used for different therapies. Finally, it is important to clearly define supportive therapy. We have found no standard definition and it would be beneficial if a standard definition for this commonly practiced intervention were to be developed.

2.2.3.2 Comparison group

Further data comparing supportive therapy both with standard care and with other treatments are needed. These should be clearly defined within the trial, even if it is routine care.

2.2.4 Outcomes

Outcomes chosen should be clinically relevant and widely used. Outcomes such as relapse or hospitalisation are less vulnerable to bias than scale data, and may therefore be preferable. Little data are available presently on adverse effects, social functioning, occupational status, quality of life, and economic outcomes, all of which would be of interest. Such outcomes are often not difficult to record over the short, medium or long term.

ACKNOWLEDGEMENTS

We are grateful to the staff of the Cochrane Schizophrenia Group for valuable support. In particular, for the original version, the search strategy was devised and run by Judith Wright, and statistical advice was provided by John Rathbone. Tessa Grant and Gill Rizzello provided administrative and copy editing support. Tor Pettit helped with protocol development, quality rating and selection of studies, data extraction and was co-author of text of the original version of this review.

For the 2012 update version Samantha Roberts ran the search, Lindsey Air and Claire Irving provided administrative and editorial support.

The Cochrane Schizophrenia Group produce and maintain a template for the methods section of their reviews. For the 2012 update we have used and adapted this.

We would like to thank M. Amr alMukhallalati, Basem Jaber, Bilal AlBaroudi and Diarmid Sinclair for their helpful observations and comments when peer reviewing the 2012 update version, and Heather Maxwell for copy editing.



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* Indicates the major publication for the study



CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Coyle 1988

Methods	Allocation: stratified random assignment. Blindness: therapists not aware of specific study hypothesis. Duration: approximately 6 months.
	Design: parallel.
	Setting: outpatient.
	Country: USA.
Participants	Diagnosis: schizophrenia. N = 94. Age: mean~47 years, range 21-72 years. Sex: 40M, 38F (data not reported for 16 participants). History: long duration of illness.
	Exclusions: a diagnosis of drug and/or alcohol abuse.
Interventions	1. Individual supportive therapeutic method: dose 10 fortnightly 90-minute sessions. N = 24.
	 Social skills training method. N = 23. Psychoeducational method. N = 23. Psychotropic medication alone. N = 24.
Outcomes	Death. Service outcomes: hospitalisation. Leaving the study early.
	Unable to use - General functioning: KAS (no SD reported).
Notes	General functioning: KAS (no SD reported).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Stratified random assignment according to age, sex and race. No further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Out of the 94 subjects entering the study, 14 did not complete the treatment program. In the individual therapy method, one subject died of medical causes, one subject was hospitalised and one subject did not complete the data;



Coyle 1988 (Continued)		in the social skills training method, three subjects dropped out and one was hospitalised; in the psychoeducational method, four subjects dropped out and two were hospitalised; in the medication only group, one subject dropped out one was hospitalised and one subject did not complete the data. In addition, there were two subjects who completed the program but a full data package	
		could not be collected on them."	
Selective reporting (reporting bias)	High risk	Reports results incompletely, no SD reported for KAS scale.	
Other bias	Unclear risk	Not enough information provided to make a judgement.	
Davidson 2004			
Methods	Allocation: randon	nised.	
	Blinding: open-label.		
	Duration: 9 months.		
	Design: parallel.		
	Setting: outpatients.		
	Country: United St		
Participants	Diagnosis: mood,	anxiety, or schizophrenia-spectrum disorder; DSM-IIIR.	
	N = 260.		
	Age: Mean 42 years (SD 11).		
	Sex: 112M, 148F		
	History: seriously ill, psychiatrically stable for the past 6 months (not hospitalised or institutionalised) and have moderate to severe impairments in social and occupational functioning.		
	Exclusions:		
Interventions		apy with consumer partner (volunteer with similar psychiatric history who has recover social network) with shared interests, within the same age range and gender of = 95.	
	the participant. N	= 95.	

- 2. Supportive therapy with non-consumer partner (volunteer without psychiatric history) with shared interests, within the same age range and gender of the participant. N = 95.
- 3. TAU: not matched with a partner. N = 70.

Participants in each treatment group were given a \$28 stipend each month.

Outcomes

Mental state: BPRS, CESD, GHQ. Global State: GAF.

Quality of Life: RSES, WBS. General Functioning: SFS.

Unable to use -

Satisfaction with service: SWS (scale not validated).

Notes



Davidson 2004 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned", no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Losses to follow-up not reported.
Selective reporting (reporting bias)	Low risk	All expected outcomes are reported.
Other bias	Low risk	The study appears to be free of other sources of bias.

Dincin 1982

Dillicili 1501			
Methods	Allocation: randomised. Blindness: none. Duration: 9 months.		
	Design: parallel.		
	Setting: outpatient.		
	Country: USA.		
Participants	Diagnosis: severely disturbed psychiatric patients (75% schizophrenia) (DSM-II). N = 132. Age: mean~25 years. Sex: 49M, 44F.		
	History: frequent referral diagnosis is schizophrenia; who are not primarily alcoholic, drug-dependent, or mentally retarded; and whose psychiatric histories include an average of three hospitalisations.		
	Exclusions: under 19 years old or attended fewer than three days during the first month after intake.		
Interventions	 Supportive treatment: dose 6 hours per week, group setting. N = 66. Comprehensive treatment: dose > 6 hours per week, full-time milieu rehabilitation programme. N = 66. 		
Outcomes	Service outcomes: hospitalisation. General functioning: gainful employment.		
	Unable to use -		



Dincin 1982 (Continued)

Service outcomes: time to hospitalisation and time spent in hospital (no SD reported).

Notes

Percentage of participants for whom no outcome data is available:

- 1. Supportive therapy 44%.
- 2. Rehabilitation programme 30%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	"Unable to contact 5 subjects from the comprehensive treatment and 14 from the supportive treatment programs. In the majority of these cases we reconstructed rehospitalisation data by talking to reliable informants and corroborating their reports with hospital records. Thus we were able to obtain verified outcome data for 50 of the 51 active subjects in the comprehensive treatment group and for 43 of the 51 active subjects in the supportive treatment group."
Selective reporting (reporting bias)	High risk	Reports results incompletely, no SDs reported for time to hospitalisation and time spent in hospital.
Other bias	Low risk	"Research was supported by grant 518 from the Illinois Department of Mental Health and Developmental Disabilities" .

Durham 2003

Methods	Allocation: randomised permuted blocking. Blindness: outcome assessors blind. Duration: 9 months treatment plus 3 months follow-up.	
Design: parallel.		
	Setting: Inpatient and outpatient.	
	Country: UK.	
Participants	Diagnosis: schizophrenia or schizoaffective disorder or delusional disorder (DSM-IV and ICD-10). N = 66. Age: mean~36 years. Sex: 45M, 21F. History: persistent and distressing hallucinations or delusions or both, stable on anti-psychotic medication for at least 6 months, duration of illness mean~13 years, range~2-31 years.	



Durham 2003 (Continued)	Exclusions: primary diagnosis of alcoholism or drug misuse, evidence of organic brain injury, history of violence,
Interventions	 Supportive psychotherapy plus treatment as usual: psychodynamic in orientation, protocol-based, delivered by staff in a community mental health team, different therapists from cognitive behavioural therapy intervention, tape recordings blindly assessed for treatment adherence. N = 23. Cognitive behavioural therapy plus treatment as usual: protocol-based, delivered by clinical nurse specialists, different therapists from supportive psychotherapy intervention, tape recordings blindly assessed for treatment adherence. N = 22. Treatment as usual. N = 21.
Outcomes	Mental state: PANSS, PSYRATS - delusions score, PSYRATS - hallucinations score. Global state: GAS. Satisfaction with treatment: patients' attitude to treatment.
	Unable to use - Global state: self-rating of degree of improvement (results not reported by group). Mental state: self-report measure of symptom severity (data not reported), self-report measure of self-esteem (data not reported), self-report measure of attitude to illness (data not reported).
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive therapy 17%. 2. CBT 5%. 3. Standard care 19%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomisation procedure (sealed envelope technique) was devised by the project statistician, and administered centrally by the non-clinical project coordinator, It was carried out separately within each treatment centre using randomised permuted blocking".
Allocation concealment (selection bias)	Low risk	"Randomisation procedure (sealed envelope technique) was devised by the project statistician, and administered centrally by the non-clinical project coordinator".
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Patients not blinded: "Patients also were asked not to mention any details of their treatment during post-treatment assessments, but three patients did." No details reported for personnel.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"Work of the independent assessors and therapists were kept strictly separate in order to maintain the blindness of the assessor".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Losses to followup or missing data balanced across intervention groups, with similar reasons for missing data. Withdrew from treatment: 1 from CBT, 4 from SPT, 2 from TAU; died from natural causes: 1 from SPT; lost to follow-up: 1 from CBT, 4 from SPT, 4 from TAU.
		"The analyses were repeated with the missing values replaced either with previous values carried forward or with group means."
Selective reporting (reporting bias)	High risk	Reports results incompletely, no data reported for self-reported symptom severity, self-esteem, and attitude to illness, results not reported by group for self-rating of degree of improvement.



Durham 2003 (Continued)

Other bias Low risk The study appears to be free of other sources of bias.

Eckman 1992

Methods	Allocation: randomised. Blindness: attempts made to keep outcome assessors blind, blindness broken in a few cases. Duration: 18 months.	
	Design: parallel.	
	Setting: not reported.	
	Country: USA.	
Participants	Diagnosis: schizophrenia (DSM-III-R). N = 41*. Age: mean~40 years. Sex: all male. History: veterans receiving constant maintenance neuroleptic drug therapy, and are able to tolerate 5-10 mg fluphenazine decanoate every other week.	
	Exclusions: evidence of mental retardation or organic brain syndrome, currently abusing alcohol or drugs.	
Interventions	 Supportive group psychotherapy: dose twice-weekly 90 minute sessions for 6 months, then onc weekly 90-minute sessions for 1 year; insight-oriented and supportive, information and education about schizophrenia provided, . N = 20. Skills training group: dose twice-weekly 90-minute sessions for 6 months, then once-weekly 90-minute sessions for a year. N = 21. 	
Outcomes	Leaving the study early.	
	Unable to use - Mental state: BPRS (data not reported by group), SANS (data not reported by group). General functioning: skill assessment by role play (SD not reported, scale not validated).	
Notes	*number randomised not reported, data reported only for participants who completed at least 6 months of psychosocial treatment and all pre- and post-treatment assessments.	
	Percentage of participants for whom no outcome data is available: 1. Supportive therapy 43%. 2. Skills training 25%.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Patients not blinded, no details reported for personnel.



Eckman 1992 (Continued)		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"Although attempts were made to keep the raters blind to the subjects' treatment conditions, a few patients revealed this information".
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	"Over the full I8 month study period, more subjects in the skills training (15/21) than in the supportive therapy (12/21) remained in the study".
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported by group for BPRS and SANS, SD not reported for skills assessment.
Other bias	Low risk	"Supported by Medical Research Service, US Department of Veterans Affairs and by NIMH grants".

Falloon 1982

-alloon 1982					
Methods	Allocation: randomised. Blindness: medication prescriber blind to treatment group, some outcome assessments blind. Duration: 2 years.				
	Design: parallel.				
	Setting: outpatients.				
	Country: USA.				
Participants	Diagnosis: schizophrenia (DSM-III). N = 39. Age: mean~26 years, range 18-41 years.				
	Sex: 26M, 13F.				
	History: definite diagnosis of schizophrenia according to PSE, residence or close daily contact with one or both biological parents, and use of English as primary language in the home.				
	Exclusions: < 18 or > 45 years old,				
Interventions	 Individual supportive therapy: dose weekly for 3 months, fortnightly for 6 months, then monthly. N = 19. Family treatment approach: dose weekly for 3 months, fortnightly for 6 months, then monthly. N = 20. 				
Outcomes	Global state: relapse, remission. Service outcomes: hospitalisation. Leaving the study early. General functioning: paid for work, admission to residential placement, admission to jail. Behaviour: SBAS, coping style. Engagement with services: poor attendance at appointments.				
	Complicance: poor compliance to therapy Medication: prescribed IM depot medication.				
	Unable to use - Service outcomes: average number of days in hospital (SDs not reported). Mental state: Hopkins' Symptom Checklist (data not reported), target symptom rating (scale not peer-reviewed), episode of depression (data not reported). General functioning: problem solving score (no SDs reported), change in work and social status (means and SDs not reported), SAS-SR (means and SDs not reported), knowledge about schizophrenia (scale not peer-reviewed).				



Falloon 1982	(Continued)
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Family coping score (no SDs reported).

Antipsychotic drug dose (no SDs reported), antipsychotic drug plasma level (data not reported), co-ef-

ficient of variation in plasma drug level/prescribed dose ratio (data not reported).

Economic outcomes: direct costs (data not reported), indirect costs (data not reported).

Notes Percentage of participants for whom no outcome data is available at 2 years:

1. Supportive psychotherapy 11%.

2. Family therapy 15%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomized", no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	"All patients were seen monthly at the clinic by a research psychiatrist or a clinical pharmacist, who was blinded to the type of treatment and responsible for managing the pharmacologic aspects of treatment." No details reported about other staff.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcomes either self-assessments or by blinded assessors. "The raters were the prescribing doctors who were blinded to the assignment to treatment."
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Two patients receiving family therapy and one receiving individual therapy withdrew from the study in the early stages of treatment". Patients who withdrew were not analysed for any outcome. Losses to follow-up or missing data balanced across intervention groups, with similar reasons for missing data.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for Hopkins' Symptom Checklist and episode of depression, SDs not reported for average number of days in hospital.
Other bias	Low risk	Funded by an NIH grant.

Haddock 1999

Methods

Blindness: outcome assessor blind to treatment allocation. Duration: 5 months treatment plus 2 years follow-up.
Design: parallel.
Setting: inpatient

Allocation: randomised.

Country: UK.

Participants Diagnosis: schizophrenia or schizoaffective disorder (DSM-IV).

N = 21.

Age: mean~29 years Sex: 19M, 2F.

History: first treatment for psychosis less than five years ago. Admitted to an acute general psychiatric ward for psychotic symptoms at time of entry into study.



Haddock 1999 (Continued)	Exclusions: not reporte	rd.
Interventions	CBT for psychosis who an inpatient stay over 5 for 4 months. N = 11. 2. Cognitive-behaviour chosis who also deliver	ng and psychoeducation: manualised, delivered by therapists with expertise in also delivered the CBT, supervised using tape recordings, therapy given during 5 weeks or shorter if the patient was discharged, then monthly booster sessions all treatment: manualised, delivered by therapists with expertise in CBT for psyred the supportive therapy, supervised using tape recordings, therapy given durver 5 weeks or shorter if the patient was discharged, then monthly booster ses-
Outcomes	Global state: relapse. Mental state: BPRS, PAI Leaving the study early Unable to use - Global state: number o Mental state: PSYRATS	f relapses per participant (SDs not reported).
	Not used in review -	
		nber of days in hospital before first discharge, total number of days in hospital ration (not used in review).
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive therapy 0%. 2. CBT 10%.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly allocated" no further details reported.

Authors' judgement Unclear risk Unclear risk	"Randomly allocated" no further details reported.
Unclear risk	
	No information reported.
Unclear risk	No information reported.
Low risk	"Independent assessment using the BPRS and PSYRATS was carried out by the project psychiatrist, who was blind to treatment allocation, and took place on entry to the study, at end of treatment and following the final booster session (4 months following discharge). Days in hospital for the initial episode and any subsequent episodes, number of relapses, time to relapse of psychotic symptoms and time to first readmission were recorded for each patient from entry to the study and for 2 years following entry to the study by a blind independent assessor using patient case notes".
Low risk	"One patient in the CBT treatment withdrew from the study after three sessions because he did not find it was helpful." Unclear if patient who withdrew imputed into outcomes.
High risk	Reports results incompletely, SDs not reported for global state: number of relapses per participant; data not reported for mental state: PSYRATS.



Haddock 1999 (Continued)

Other bias Unclear risk Not enough information provided to make a judgement.

Hogarty 1997-study 1

Methods	Allocation: randomised. Blindness: none. Duration: 3 years.
	Design: parallel.
	Setting: outpatients.
	Country: USA.
Participants	Diagnosis: schizophrenia or schizoaffective disorder (RDC). N = 97. Age: mean~29 years, range 16-55 years. Sex: 56M, 41F.
	History: living with family, IQ above 75, absence of organic brain syndrome and serious alcohol or drug abuse or dependence in the previous 6 months, recruited whilst in hospital, treated in the community.
	Exclusions: Age $<$ 16, or $>$ 55; medical contraindications that precluded taking maintenance antipsychotic medication.
Interventions	 Supportive therapy: biweekly, provided by the same therapists as personal therapy or family therapy, manualised, supervised. N = 24. Personal therapy: CBT, 3-stage approach, sought to enhance personal and social adjustment through the identification and effective management of affect dysregulation, manualised, weekly but with less contact in year 3 for those who completed treatment objectives, provided by therapists who also provided supportive therapy, but not family therapy. N = 23. Family therapy: manualised, biweekly, provided by therapists who also provided supportive therapy but not personal therapy. N = 24. Personal therapy plus family therapy. N = 26.
Outcomes	Global state: relapse. Mental state: episode of affective symptoms. Leaving the study early: treatment-related reasons.
	Unable to use - Global state: GAS, Subjective Response Questionnaire (data not reported). Mental state: BPRS, Raskin Depression Scale, Covi Anxiety Scale, Wing Negative Symptom Scale, Everyday Worries Scale (data not reported). General functioning: KAS, SAS II, Major Role Adjustment Inventory (data not reported). Satisfaction with treatment (data not reported by group).
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive psychotherapy 33%. 2. CBT 4%. 3. Family therapy 21%. 4. CBT plus family therapy 4%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" no further details reported.



Hogarty 1997-study 1 (Continu	ued)	
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	High risk	"The ratings of treating clinicians who were not blind to the treatment conditions, thereby raising the question of rater bias", "results should be viewed cautiously in the absence of independent and blind clinical assessors."
Incomplete outcome data (attrition bias) All outcomes	Low risk	8 patients in the supportive therapy group had treatment-related terminations, one in the personal therapy group, five in the family therapy group and 1 in the combined personal and family therapy group.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for Global state: GAS, Subjective Response Questionnaire; Mental state: BPRS, Raskin Depression Scale, Covi Anxiety Scale, Wing Negative Symptom Scale, Everyday Worries Scale; General functioning: KAS, SAS II, Major Role Adjustment Inventory; data not reported by group for Satisfaction with treatment.
Other bias	Low risk	"Supported by a MERIT extension of NIMH grant."

Hogarty 1997-study 2

Methods	Allocation: randomised. Blindness: none. Duration: 3 years.
	Design: parallel.
	Setting: outpatient.
	Country: USA.
Participants	Diagnosis: schizophrenia or schizoaffective disorder (RDC).
	N = 54. Age: mean~33 years, SD~8 years, range 16-55 years.
	Sex: 24M, 30F.
	History: living independent of family, IQ above 75, absence of organic brain syndrome and serious al- cohol or drug abuse or dependence in the previous 6 months, recruited whilst in hospital, treated in the community.
	Exclusions: Age<16, >55; medical contraindications that precluded taking maintenance antipsychotic medication.
Interventions	1. Supportive therapy: biweekly, provided by the same therapists as personal therapy or family therapy, manualised, supervised. N = 29.
	2. Personal therapy: CBT, 3-stage approach, sought to enhance personal and social adjustment through the identification and effective management of affect dysregulation, manualised, weekly but with less contact in year 3 for those who completed treatment objectives, provided by therapists who also provided supportive therapy, but not family therapy. N = 25.
Outcomes	Global state: relapse. Mental state: episode of affective symptoms.
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Hogarty 1997-study 2 (Continued)

Unable to use -

Global state: GAS, Subjective Response Questionnaire (data not reported).

Mental state: BPRS, Raskin Depression Scale, Covi Anxiety Scale, Wing Negative Symptom Scale, Every-

day Worries Scale (data not reported).

General functioning: KAS, SAS II, Major Role Adjustment Inventory (data not reported).

Satisfaction with treatment (data not reported by group).

Notes Percentage of participants for whom no outcome data is available:

1. Supportive psychotherapy 17%.

2. CBT 16%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	High risk	"The ratings of treating clinicians who were not blind to the treatment conditions, thereby raising the question of rater bias", "results should be viewed cautiously in the absence of independent and blind clinical assessors."
Incomplete outcome data (attrition bias) All outcomes	Low risk	Five patients in the supportive therapy group and four in the personal therapy group had treatment-related terminations.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for Global state: GAS, Subjective Response Questionnaire; Mental state: BPRS, Raskin Depression Scale, Covi Anxiety Scale, Wing Negative Symptom Scale, Everyday Worries Scale; General functioning: KAS, SAS II, Major Role Adjustment Inventory; data not reported by group for Satisfaction with treatment.
Other bias	Low risk	"Supported by a MERIT extension of NIMH grant."

Kemp 1996

Methods	Allocation: randomised. Blindness: some outcome assessments blind. Duration: 18 months follow-up.
	Design: parallel.
	Setting: outpatients.
	Country: UK.
Participants	Diagnosis: schizophrenia or severe affective disorder or schizophreniform disorder or schizoaffective disorder or delusional disorder or psychotic disorder not otherwise classified (DSM-III-R). N = 74.



Kemp 1996 (Continued)	
	Age: mean~36 years. Sex: 39M, 35F. History: no people with low IQ, no deafness, no organic brain disease, no non-English speakers, hospitalised.
Interventions	 Supportive counselling: dose 4-6 sessions lasting 20-60 minutes twice a week plus 3 booster sessions, counselling non-specific, concerns listened to but therapists declined to discuss treatment. N = 35. Compliance therapy: dose 4-6 sessions plus 3 booster sessions of cognitive behavioural and motivational interviewing techniques. N = 39.
Outcomes	Service outcomes: rehospitalisation. Mental state: BPRS. Leaving the study early. Global state: GAF. Attitude to medication: AMQ, DAI Insight: SAI. Economic outcomes: direct costs. Unable to use - Death (data not reported by group). Adverse effects: Simpson-Angus scale for extra-pyramidal side effects, Barnes akathisia scale (data not reported). Medication compliance measure (scale not peer reviewed). NART- National Adult Reading Test (data not reported).
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive therapy 43%. 2. CBT 28%.
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomly assigned by means of a table of random numbers to compliance therapy or control treatment"
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"Initial compliance was rated blind to intervention by the patients' primary nurses,"; "The evaluations at six months were carried out by an independent assessor (a community psychiatric nurse) blind to intervention."; "Most of the ratings of functioning and compliance initially and at three months were made by a research psychiatrist, who was not blind to intervention group. However, the compliance ratings were based on information from impartial sources, including community psychiatric nurses and outpatient doctors on the clinical teams. By contrast, the ratings at six months were carried out by a researcher trained in the use of all the ratings, who was blind to intervention group." Some outcome assessors blinded, except third month assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Overall, 35% were lost during the follow-up period (11 (28%) from the compliance therapy group and 15 from. the control group (43%); Ten participants were lost during the first six months of follow-up, five from each group; Subsequently, a further 14 could not be directly re-interviewed at 12-month fol-



Kemp 1996 (Continued)		low-up, Post-treatment drop-out occurred due to refusal (11), patients either being uncontactable (7) or moving out of the area (2) or recovered patients who were working or at college and no longer having psychiatric out-patient treatment (2). One patient was in prison, There were two deaths, one resulting from stroke, and one from myocardial infarction". Details reported for imputing missing data by method of maximum likelihood.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for adverse effects: Simpson-Angus scale for extra-pyramidal side effects, Barnes akathisia scale, NART; data not reported by group for death; scale not peer reviewed for medication compliance measure.
Other bias	Low risk	Funding from the Medical Research Council.

Klein 1998			
Methods	Allocation: randomised.		
	Blinding: open label.		
	Duration: 6 months.		
	Design: parallel.		
	Setting: outpatients.		
	Country: United States.		
Participants	Diagnosis: severe mental illness (DSM IV criteria for schizophrenia/ chronic major mood disorder) and substance abuse.		
	N = 61.		
	Age: 26-59 years; mean 40 years.		
	Sex: 45M, 16F.		
	History: severe mentally ill patients stable enough to be in the community for the duration of the study period and relationally connected to their Intense Case Manager (ICM), previously admitted to mental hospital or inpatient psychiatric unit.		
	Exclusions: previously been assigned a Friend's Connection (FC) or presently working with an FC.		
Interventions	1.Peer support plus intensive case management: at least 2-3 times/week meetings with Friend's Connection, a supplemental rehabilitation intervention that provides one-to-one support including friendship, counselling, social support, and meaningful leisure time activities. N = 10.		
	2. Intensive case management plus treatment as usual. N = 51.		
Outcomes	Leaving the study early.		
	Global state: Relapse (defined as a "crisis event": requiring emergency room services (medical or psychiatric), threatening staff members or other individuals, making suicide attempts or threats, creating a public disturbance, and disappearing from the place of residence for more than 1 day).		
	Costs of inpatient care.		
	Unable to use:		
	Service Outcomes: hospitalisation - average inpatient days (no SDs).		



Kle	in 1	L998	(Continued)
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Quality of life: Lehman's Quality of Life Questionnaire (no SDs).

Global state: GAF, average social interaction per week (no SDs). Activities of daily living - mean frequency of activities (no SDs).

Costs - use of community resources (no data).

Notes

2 alternates were randomly selected for each of the 10 clients. 9/10 in the study sample were alternates. The 9 clients originally selected for the sample refused to participate and were included in the comparison group.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"The study sample was randomly selected from ICM caseload listsThe remaining 51 clients in the study population served as a comparison group", no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	"One participant who originally consented to the study and was assigned an FC withdrew after a few weeks. We did not replace her as she was already part of the study group, and all analyses include her".
Selective reporting (reporting bias)	High risk	The study did not report SDs for hospitalisation, quality of life, activities of daily living and social functioning. No data were reported for use of community resources.
Other bias	Low risk	Appears free of other sources of bias.

Levine 1998

Methods	Allocation: randomised. Blindness: outcome assessment blind. Duration: 6 weeks treatment plus 4 weeks follow-up.
	Design: parallel.
	Setting: not reported.
	Country: Israel.
Participants	Diagnosis: schizophrenia (DSM-III-R). N = 12. Age: mean~35 years, range 24-42 years. Sex: not reported.



Levine 1998 (Continued)	History: mean duration of illness~11 years, range 6-20 years, 8 years of schooling or more, active delusional system, no change of antipsychotic drugs given in standard doses in the last 3 months
	Exclusions: alcohol and/or drug abuse, chronic physical condition, orthodox (Jewish) religious conviction.
Interventions	 Supportive group: weekly for 6 weeks plus one follow-up session, with avoidance of relating the discussion to delusional experiences. N = 6. Cognitive-dissonance group therapy: dose weekly for 6 weeks plus one follow-up session; modified cognitive therapy. N = 6.
Outcomes	Mental state: PANSS.
Notes	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" . No further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"The PANSS scale was scored by an independent, board certified, senior psychiatrist well trained in the use of the PANSS. The psychiatrist was not aware of the group assignment of the subjects." No further details reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Losses to follow-up not reported.
Selective reporting (reporting bias)	High risk	Not all expected outcomes are reported.
Other bias	Unclear risk	Not enough information provided to make a judgement.

wis 2002h

Lewis 2002b	
Methods	Allocation: randomised with stratification. Blindness: outcome assessment blind. Duration: 5 weeks therapy plus 5 weeks follow-up.
	Design: parallel.
	Setting: inpatients & outpatients.
	Country: UK.
Participants	Diagnosis: schizophrenia or schizoaffective disorder or delusional disorder (DSM-IV). N = 315*.



Lewis 2002b (Continued)

Age: median~27 years.

Sex: 216M, 93F.

History: inpatients or day patients, 83% in first admission and 17% in second admission, positive psychotic symptoms for 4 weeks or more, moderate or severe score on PANSS for delusions or hallucinations, neither substance misuse nor organic disorder judged to be the major cause of psychotic symptoms.

Exclusions: Substance misuse or organic disorder judged to be the major cause of psychotic symptoms. Patients legally detained in hospital.

Interventions

- 1. Supportive counselling plus routine care: 15-20 hours of therapy within 5 weeks plus 3 booster sessions; manualised, supervised, audio taped sessions blindly rated for treatment fidelity and quality, provided by therapists trained in CBT in psychosis who also provided the CBT intervention. N = 106. 2. Cognitive- behavioural therapy plus routine care: dose 15-20 hours of therapy within 5 weeks plus 3 booster sessions; manualised, supervised, audio taped sessions blindly rated for treatment fidelity and quality, provided by therapists trained in CBT in psychosis who also provided the supportive counselling intervention. N = 101.
- 3. Routine care: not standardised. N = 102.

Outcomes

Mental state: PANSS, PSYRATS. Leaving the study early.

Notes

*Seven participants excluded after randomisation.

Percentage of participants for whom no outcome data is available:

- 1. Supportive psychotherapy 17%.
- 2. CBT 5%.
- 3. Treatment as usual 19%.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Independent, concealed, randomisation of individuals with minimisation was then performed by trial administrator at each centre. Stratification was undertaken with variables".
Allocation concealment (selection bias)	Low risk	"independent, concealed, randomisation of individuals with minimisation was then performed by trial administrator at each centre. Stratification was undertaken with variables"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	"Interventions carried out independently of clinical staff who were kept unaware of treatment allocation."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"All outcome assessments were made blind to treatment allocation. Extensive steps were taken to maintain blindness of raters. In all cases randomisation was carried out by a trial administrator independently of rater or therapist. Therapist and rater were not to communicate details about individual patients to each other. Office space and data storage were kept separate and secure. Clinical staff were instructed not to divulge details of therapist contacts to the raters."
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Fifteen patients (CBT 4; supportive counselling 4; routine care 7) withdrew consent to participate during the follow-up period, but are included in the analysis prior to their withdrawal; 13 of these withdrawals occurred during the first 2 weeks. One patient died during the follow-up period (in the supportive counselling group). "Analyses were conducted on an intention-to-treat



Lewis 2002b (Continued)		basis, with patients analysed in the treatment group to which they were randomised"
Selective reporting (reporting bias)	Low risk	All expected outcomes are reported
Other bias	Low risk	The trial was funded as follows: UK Medical Research Council (41%): Northwest England NHSE Office (27%); Trent NHSE Office (7%); the following health authorities: Manchester (8%); Salford and Trafford (2%); Liverpool (3%); Sefton (3%); St Helens and Knowsley (3%); North Nottinghamshire (6%).

Malm 1982

Malm 1982	
Methods	Allocation: randomised. Blindness: none. Duration: 12 months treatment plus 12 months follow-up.
	Design: parallel.
	Setting: inpatients & outpatients.
	Country: Sweden.
Participants	Diagnosis: schizophrenic psychosis. N = 80. Age: 18-50 years. Sex: 43M, 25F. History: total period of hospitalisation less than 3 years, no organic brain damage, no serious somatic illness, IQ > 70.
	Exclusions: Gross organic brain damage, serious somatic illness, epilepsy or pathological EEG (which motivated treatment with antiepileptic drugs), narcotic drug addictions, dependence on alcohol, known sex chromosome aberration, mental retardation (IQ < 701, and previous social skills training or group therapy, patients who recovered after the first month of treatment with neuroleptics alone.
Interventions	 Group therapy plus fluphenazine depot plus social skills training: dose weekly sessions of 60-90 minutes, group therapy lasting 1 year; therapists not formally trained, manualised and supervised, communication-oriented. N = 40. Fluphenazine depot plus social skills training: dose 10 hours per week of individual and group training. N = 40.
Outcomes	Death. Global state: relapse, no remission. Service outcomes: not discharged from hospital. Mental state: CPRS - inability to feel and derealisation items. Leaving the study early. General functioning: KAS - free time activities subscale and withdrawal syndrome. Unable to use - Mental state: CPRS - global rating of illness, all items except derealisation and inability to feel (data not reported). General functioning: KAS - socially expected activities subscale, offensive behaviour syndrome and
	work and study item (data not reported), life events (data not reported).
Notes	Percentage of participants for whom no outcome data is available: 1. Group therapy plus fluphenazine depot plus social skills training 15%. 2. Fluphenazine depot plus social skills training 15%



Malm 1982 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"By randomisation they were allocated to the therapy or the control group.", stratified randomisation based on sex.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"Ratings agreed closely with those of the independent blind raters." Raters that were blinded not specified, "There were 7 independent raters, 5 physicians and 2 psychologists."
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Six patients in the therapy group and six in the control group either did not start or did not complete the study" Details of dropouts further reported in table. Dropouts not included in analysis.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for CPRS - global rating of illness, all items except derealisation and inability to feel; KAS - socially expected activities subscale, offensive behaviour syndrome and work and study item
Other bias	Low risk	The study appears to be free of other sources of bias. The study has been supported by grants from the Einar Hansen Fund for Research, the Leo Research Fund, the Lundbeck Foundation and the Stiftelsen Psykiatriska Forskningsfonden, Goteborg.

O'Donnell 1999

O Donnett 1999	
Methods	Allocation: randomised. Blindness: none. Duration: 12 months.
	Design: parallel.
	Setting: inpatients & outpatients.
	Country: Australia.
Participants	Diagnosis: schizophrenia (79) or schizoaffective disorder (19) or schizophreniform disorder (7) or bipolar affective disorder (14) (DSM-IV). N = 119. Age: mean~36 years. Sex: 68M, 51F. History: English-speaking, mean duration of illness~10 years, referred for case management by community health services.
	Exclusions: Age <18 or >65 years, co-diagnosis substance dependence disorder, no significant history of violence unless associated with acute psychosis, no extant community treatment order or community counselling order.



O'Donnell 1999 (Continued)

Interventions

- 1. Advocacy plus client-focused case management: advocacy provided by people who had experienced mental illness themselves, siblings or carers of people who had experienced mental illness and non-consumers interested in working with people with mental illness. N = 45.
- 2. Client-focused case management. N = 39.
- 3. Standard case management. N = 35.

Outcomes

Death.

Leaving the study early.

Unable to use -

Global state: HoNOS, GAF(data not reported).

Service outcomes: hospitalisation, duration of hospital admissions, use of crisis services (data not re-

ported).

Mental state: response to treatment (data not reported).

General functioning: LSP (data not reported).

Satisfaction with treatment: CSQ (data not reported).

Quality of life: Quality of Life Index for Mental Health (modified) (data not reported).

Family Interview Schedule (data not reported).

Compliance (data not reported).

Notes

Percentage of participants for whom no outcome data is available:

- 1. Supportive psychotherapy plus client-focused case management 38%.
- 2. Client-focused case management 18%.
- 3. Standard case management 43%.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly allocated" . No further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	15 participants from the standard case management group left the study early, 7 from the client-focused case management group, 18 from the client-focused case management plus advocate group. Reasons similar across groups.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for global state: HoNOS, general functioning; service outcomes: hospitalisation, duration of hospital admissions, use of crisis services, mental state: response to treatment; satisfaction with treatment; quality of life; Family Interview Schedule and compliance.
Other bias	Low risk	"Commonwealth Innovative Grants Program of the National Mental Health Strategy (Grant No. 22004) for funding this study"



Penn 2009			
Methods	Allocation: randomised.		
	Blinding: single.		
	Duration: 12 weeks.		
	Design: parallel.		
	Setting: outpatients.		
	Country: United States.		
Participants	Diagnosis: schizophrenia or schizoaffective disorder; DSM IV criteria.		
	N = 65.		
	Age: 18-65.		
	Sex: 33M, 32F.		
	History: 1)diagnosis of schizophrenia or schizoaffective disorder; 2) age 18-65 years; 3) presence of current persistent auditory hallucinations of at least moderate severity (i.e.		
	had a rating of at least 4 on the Positive and Negative Syndrome Scale) (PANSS); 4) have undergone at least two pharmacological trials, one of which being an atypical neuroleptic or clozapine for 8 weeks prior to randomisation.		
	Exclusions: 1) mental retardation (based on both IQ and functional impairment criteria) 2) current substance dependence.		
Interventions	1. Cognitive-behavioural therapy (CBT). N = 32.		
	2. Enhanced supportive therapy (ST). N = 33.		
Outcomes	Leaving the study early.		
	Service use: hospitalisation.		
	Mental state: BVAQ-R, BDI-II, PSYRATS voices scale.		
	General functioning: SFS.		
	Insight: BCIS.		
	Quality of life: RSES.		
	Not used in review -		
	Average number of days hospitalised.		
Notes			

Notes

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomisation was stratified by gender to ensure equal numbers across groups using a computer randomisation generator".
Allocation concealment (selection bias)	Low risk	"Randomisation to treatment condition (with condition being designated by a random number), was conducted by two research assistants who were blind to the correspondence between random number and treatment group."



Penn 2009 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Participants not blinded. No information reported about personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Allresearch evaluations were conducted by two research assistants (RAs) who were blind to treatment assignment. Blindness was maintained by asking participants not to talk to the RAs about their treatment. In addition, the RAs had minimal contact with the study therapists. Finally, RAs were kept blind to the coding system used to denote group membership." "Participants were assessedby research assistants blind to treatment group."
Incomplete outcome data (attrition bias) All outcomes	Low risk	"We analysed data on all participants irrespective of treatment adherence (i.e., intent-to-treat analyses) and number of post treatment assessments".
Selective reporting (reporting bias)	Low risk	All expected outcomes are reported.
Other bias	Low risk	Appears free of other sources of bias.

Pinto 1999

Methods	Allocation: randomised. Blindness: none. Duration: 6 months.		
	Design: parallel.		
	Setting: inpatients & outpatients.		
	Country: Italy.		
Participants	Diagnosis: schizophrenia (DSM-IV).		
	N = 41. Age: mean~35 years, SD~11 years.		
	Sex: 28M, 13F.		
	History: treatment-refractory psychosis, recently started clozapine, illness duration mean 12 years.		
	Exclusions: evidence of current substance abuse or organic pathology.		
Interventions	1. Supportive therapy: individual, same duration and frequency of sessions as CBT plus social skills training. N = 21.		
	2. Cognitive-behavioural therapy plus social skills training. N = 20.		
Outcomes	Global state: relapse.		
	Service outcomes: hospitalisation.		
	Mental state: BPRS, SAPS, SANS.		
	Leaving the study early.		
	Unable to use -		
	Dose of antipsychotic medication (data not reported).		
Notes	Percentage of participants for whom no outcome data is available:		
	1. Supportive psychotherapy 14%.		
	2. CBT 5%.		



Pinto 1999 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned " No further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	"One subject in the cognitive-behavioral group did not complete the study; the client developed leukopenia, and clozapine was discontinued. In the supportive therapy group, two subjects withdrew because of refusal to participate further, and a third developed seizures and his clozapine was stopped." Losses to followup not included in analysis.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for dose of antipsychotic medication.
Other bias	Unclear risk	Not enough information provided to make a judgement.

Sensky 2000b

Methods	Allocation: randomised. Blindness: outcome assessments blind. Duration: 9 months treatment plus 9 months follow-up.		
	Design: parallel.		
	Setting: not reported.		
	Country: UK.		
Participants	Diagnosis: schizophrenia (ICD-10 and DSM-IV). N = 90. Age: mean~39 years, 95% CI 37-42 years. Sex: 53M, 37F. History: mean length of illness~14 years, 95% CI 12-17 years, persistent symptoms resistant to medication, complaining of positive symptoms or depression. Exclusions: primary diagnosis of alcohol or drug abuse, current abuse of drugs or alcohol warranting specific clinical intervention such as attendance at a specialist substance misuse clinic, exclusively negative symptoms or not complaining of any positive symptoms or of depression.		
Interventions	1. Befriending: dose mean number of sessions~19; provided by therapists trained in CBT, provided by the same therapists as CBT, sessions audiotape for supervision and quality control, psychotic or affective symptoms not directly tackled. N = 44.		



Sensky 2000b (Continued)	2. Cognitive-behavioural therapy: dose mean number of sessions~19; manualised, provided by therapists trained in CBT, provided by the same therapists as befriending, sessions audiotape for supervision and quality control. N = 46.
Outcomes	Mental state: CPRS, MADRS, SANS. Leaving the study early. Medication: no reduction in dose of antipsychotic medication. Unable to use - Satisfaction with treatment: patient satisfaction score (scale not peer reviewed).
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive psychotherapy 14%. 2. CBT 20%.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Simple randomisation applied". No further details reported.
Allocation concealment (selection bias)	Low risk	"The randomisation was by members of the research team not involved with either the assessments or the treatments."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Assesors were independent of the randomisation process and remained blind to each patient's assigned group throughout the study".
Incomplete outcome data (attrition bias) All outcomes	Low risk	Missing data have been imputed using appropriate methods "intention to treat analysis".
Selective reporting (reporting bias)	Low risk	Patient satisfaction score scale not peer reviewed.
Other bias	Low risk	Funded by grant from Welcome trust, further financial support was provided by Hounslow and Spelthorne community and mental health service trust.

Spaulding 1999

Methods	Allocation: matched pair randomisation. Blindness: some outcomes assessments blind. Duration: 6 months treatment plus 1 year of continued standard rehabilitation.
	Design: parallel.
	Setting: inpatient.
	Country: USA.
Participants	Diagnosis: schizophrenia (DSM-III-R).



Spaulding 1999 (Continued)

N = 91.

Age: mean~36 years, SD~10 years.

Sex: 56M, 35F.

History: hospitalised, failure in all other available treatment settings, not responding to short-term inpatient

treatment sufficiently enough to allow discharge, having a primary chart diagnosis of an Axis I psychiatric

disorder, IQ > 70.

Exclusions: primary diagnosis of mental retardation or substance abuse, no dangerous behaviour requiring a higher security setting.

Interventions

1. Supportive group therapy plus standard regimen: dose 3 sessions per week lasting 45-60 minutes; manualised, videotaped sessions assessed using Q-sort instrument, therapists experienced in supportive group therapy, standard regimen consists of comprehensive psychiatric rehabilitation. N = 42.

2. Cognitive treatment: dose 3 sessions per week lasting 45-60 minutes cognitive sub programmes of Integrated Psychological Therapy, group, manualised, therapist trained in Integrated Psychological Therapy, standard regimen consists of comprehensive psychiatric rehabilitation. N = 49.

Outcomes

Leaving the study early.

Unable to use -

Mental state: BPRS, PANSS, TLC, NOSIE-30 (data not reported).

General functioning: AIPSS, Symptom Management skill assessment, Medication Management skill assessment, Leisure Skills assessment, Conversational Skills assessment, COGLAB cognitive assessment battery, Trailmaking test, Tactile Performance Test, Categories, Rey Auditory and Visual Learning Tasks, Denman Neuropsychological Memory Test (data not reported).

Notes

Percentage of participants for whom no outcome data is available:

- 1. Supportive psychotherapy 0%.
- 2. CBT 2%.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Patients in the cohort were matched pair-wise, as closely as possible, for gender, chronicity, and overall levels of personal and social functioning. Each patient in the pair was then assigned to one of two groups. When there was an odd number of patients in the cohort, the unmatched individual was randomly assigned to one of the groups. One of the groups was then randomly assigned to the experimental condition and the other to the control condition". No further details reported.
Allocation concealment (selection bias)	Unclear risk	Matched pair randomisation. No further details reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	"All clinical and research personnel were blind to experimental condition, except for the cognitive and supportive therapists, the observers who performed the process measures in the project's first year, and the principal investigator."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"All NOSIE-30 raters were blind to treatment group assignment"; "There definitely were no breaches with any staff directly involved in clinical or laboratory assessments". "All clinical and research personnel were blind to experimental condition, except for the cognitive and supportive therapists, the observers who performed the process measures in the project's first year, and the principal investigator."



Spaulding 1999 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	"One subject who dropped out after the intensive treatment period began."
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for BPRS, PANSS, TLC, NOSIE-30, AIPSS, Symptom Management skill assessment, Medication Management skill assessment, Leisure Skills assessment, Conversational Skills assessment, COGLAB cognitive assessment battery, Trailmaking test, Tactile Performance Test, Categories, Rey Auditory and Visual Learning Tasks, Denman Neuropsychological Memory Test
Other bias	Low risk	Funded by an NIH grant

Stanton 1984

Methods	Allocation: randomised. Blindness: some outcome assessments blind, attempts made to blind therapists to hypothesis and outcomes. Duration: 2 years, had to stay in therapy for 6 months to be included in 2 year follow-up.
	Design: parallel.
	Setting: inpatient.
	Country: USA.
Participants	Diagnosis: schizophrenia or probable schizophrenia (Psychiatric Status Schedule and WHO Discrimination Criteria for Schizophrenia). N = 164.
	Age: mean 22 years.
	Sex: 112M, 52F. History: minimal prior treatment, can function outside of a hospital in some major role, without medications for 4 consecutive months preceding 2 years.
	Exclusions: history of alcohol dependency or drug use, obvious organic impairments.
Interventions	 Reality-adaptive supportive psychotherapy (RAS): dose up to once weekly. N = 76. Exploratory insight-oriented psychotherapy (EIO): dose three or more times per week of analytic psychotherapy. N = 88.
Outcomes	Leaving the study early.
	Unable to use - Service outcomes: rehospitalisation, number of days hospitalised (> 50% lost to follow-up). Mental state: Camarillo Dynamic Assessment Scale, IMPS, Psychotherapy Outcome Interview (> 50% lost to follow-up). General functioning: number of days dependent, units of productive activity, days employed full-time, occupational level reached, self-support, household responsibilities, significant relationships (> 50% lost to follow-up). Number of job changes (> 50% lost to follow-up). Days on antipsychotic medication (> 50% lost to follow-up). Visual-Verbal Test - total misses (> 50% lost to follow-up). Global state: Menninger Health-Sickness Rating Scale, Psychotherapy Outcome Interview (data not reported). Mental state: Psychiatric Status Schedule, IMPS, Inpatient Multidimensional Psychiatric Scale, Camarillo Dynamic Assessment Scale (data not reported). General functioning: WAIS, KAS, Gottschalk Social Alienation/Personal Disorganisation Scale (data not reported).



Stanton:	1984	(Continued)
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Rorschach test, Thematic Apperception Test, Visual-Verbal Test, Soskis Attitude Toward Illness Questionnaire (data not reported).

Notes

Percentage of participants for whom no outcome data is available at 6 months:

- 1. Supportive psychotherapy 32%.
- 2. Psychodynamic therapy 51%.

Percentage of participants for whom no outcome data is available at 12 months:

- 1. Supportive psychotherapy 51%.
- 2. Psychodynamic therapy 75%.

Percentage of participants for whom no outcome data is available at 24 months:

- 1. Supportive psychotherapy 67%.
- 2. Psychodynamic therapy 75%.
- > 50% of participants lost to follow-up at 12 months and 24 months, therefore data excluded.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Assignment to EIO or RAS was achieved by a random numbering system within an unrelated office."; "On those occasions where imbalances appeared, a weighting was introduced into the randomisation procedure to reestablish comparability of the EIO and RAS groups at baseline."
Allocation concealment (selection bias)	Unclear risk	"Assignment to EIO or RAS was achieved by a random numbering system within an unrelated office" Details of method not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	"We knew that significant others, ward staff, and project staff also would not be blind to the patient's experimental condition."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	"Four different groups of people administered the measures: (1) project psychiatrists; (2) blind evaluators; (3) psychological testers; and (4) research assistants."; "Psychological testers were clinical psychologists with expertise in psychological testing. They were blind to the experimental conditions to which patients were assigned and also unfamiliar with the study design and hypotheses."
Incomplete outcome data (attrition bias) All outcomes	High risk	High attrition rates reported throughout both groups during all three follow-up periods (See Notes above). Reasons for attrition not reported.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for Menninger Health-Sickness Rating Scale, Psychotherapy Outcome Interview, Psychiatric Status Schedule, IMPS, Inpatient Multidimensional Psychiatric Scale, Camarillo Dynamic Assessment Scale, WAIS, KAS, Gottschalk Social Alienation/ Personal Disorganisation Scale, Rorschach test, Thematic Apperception Test, Visual-Verbal Test, Soskis Attitude Toward Illness Questionnaire;
Other bias	Low risk	"supported in part by NIMH Grant "

Tarrier 1998

Methods Allocation: stratified block randomisation.

Blindness: outcome assessments blind.

Duration: 3 months treatment plus 2 years follow-up.



Tarrier 1998 (C	Continued	1)
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Design: parallel.

Setting: outpatient.

Country: UK.

Participants

Diagnosis: schizophrenia or schizoaffective psychosis or delusional disorder (DSM-III-R).

N = 87

Age: mean~39 years. Sex: 69M, 18F.

History: median length of illness~11 years, range 1-42 years, persistent hallucinations or delusions or both for a minimum of 6 months and at least 1 month of stabilisation if they had experienced an exacerbation within this period, stable medication.

Exclusions: organic brain disease, substance abuse as primary problem, threat of violence towards assessors, psychological or family intervention.

Interventions

- 1. Supportive counselling plus routine care: dose 20 sessions of 1 hour twice a week for 3 months plus monthly booster sessions for 4 months. N = 26.
- 2. Intensive cognitive behaviour therapy plus routine care: dose 20 sessions of 1 hour twice a week for 3 months plus monthly booster sessions for 4 months; coping strategy enhancement, problem solving and relapse reduction strategies used. N=33.
- 3. Routine care: standard psychiatric management with medication and monitoring outpatient follow-up and the care programme approach. N = 28.

Outcomes

Death.

Global state: relapse.

Mental state: clinically important improvement in general mental state, SANS, BPRS based positive

symptoms score. Leaving the study early.

Unable to use -

Mental state: BDI, BHS, KGV (data not reported), PSE (SDs not reported)

General functioning: SFS (data not reported).

Dose of antipsychotic medication (data not reported).

Notes

Percentage of participants for whom no outcome data is available:

- 1. Supportive therapy 4%.
- 2. CBT 12%.
- 3. Standard care 4%.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Patients were allocated on a stratified block randomised procedure with block size equal to nine by using severity of symptoms and sex to one of three treatment groups"
Allocation concealment (selection bias)	Unclear risk	"Allocation, contained in sealed envelopes, was carried out independently by a third party."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Effort was made to blind the independent assessors to treatment allocation by using separate offices in a different part of the hospital for the assessors and therapists, using separate administrative procedures, instructing patients



Tarrier 1998 (Continued)		
		to not reveal details of their care, data entry being carried out independently of the assessors, and using a multiple coding system for treatment groups."
Incomplete outcome data (attrition bias) All outcomes	Low risk	"Analysis by intention to treat was completed on the 87 allocated patients, with last observations (scores before treatment) being carried forward for the eight patients for whom scores after treatment were missing". Missing data have been imputed using appropriate methods (intention to treat analysis), and losses to follow-up or missing data balanced across intervention groups.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for BDI, BHS, KGV, SFS, and dose of antipsychotic medication; SDs not reported for PSE.
Other bias	Low risk	Funded by a Wellcome Trust grant.

Turkington 2000

Risk of bias	
Notes	Percentage of participants for whom no outcome data is available: 1. Supportive psychotherapy 0%. 2. CBT 20%.
Outcomes	Leaving the study early. Unable to use - Service outcomes: time spent in hospital (SDs not reported). Mental state: CPRS, MADRS (SDs not reported).
Outcomes	Leaving the study early
Interventions	 Befriending: dose 6 sessions over 2 months; structured and supportive, some sessions audio taped and analysed for treatment fidelity. N = 6. Cognitive techniques: dose 6 sessions over 2 months; manualised, some sessions audio taped and assessed for treatment fidelity. N = 13.
	Exclusions: not reported.
	N = 19. Age: mean~41 years. Sex: 9M, 9F. History: mean duration of illness for befriending group~13 years, mean duration of illness for cognitive therapy group~9.2 years.
Participants	Diagnosis: schizophrenia (ICD-10).
	Country: UK.
	Setting: inpatient & outpatient.
	Design: parallel.
Methods	Allocation: randomised. Blindness: patients, assessors and clinical teams blind to allocation. Duration: 2 months therapy plus 4 months follow-up.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Patients randomised on 2:1 basis after initial assessments made. No further details reported.



Turkington 2000 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Details of method not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	"Patients randomly allocated to two groupswith the patients, assessors, and clinical team remaining blind to the allocation."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Patients randomly allocated to two groupswith the patients, assessors, and clinical team remaining blind to the allocation.
Incomplete outcome data (attrition bias) All outcomes	Low risk	"One patient in the cognitive techniques group withdrew consent after the initial interview, and so had to be excluded from the study", "Calculations were also made excluding his results, from the analysis"
Selective reporting (reporting bias)	High risk	Reports results incompletely, SDs not reported for time spent in hospital, CPRS, MADRS.
Other bias	Low risk	The study appears to be free of other sources of bias.

Uzenoff 2007

Methods	Allocation: randomised.
	Blinding: unclear.
	Duration: 6 months.
	Design: parallel.
	Setting: outpatients.
	Country: USA.
Participants	Diagnosis: schizophrenia, schizoaffective disorder or schizophreniform. DSM IV criteria.
	N = 24.
	Age: ≥ 16.
	Sex: 14 M, 10F
	History: clinically stable outpatients recovering from a first psychotic episode (in treatment <12 months).
	Exclusions: not reported.
Interventions	1. Adherence-Coping Education (ACE). N = 13.
	2. Supportive Therapy (ST). N = 11.
Outcomes	Leaving the study early.
	Medication: adherence, ROMI (adherence and nonadherence).
	Need for treatment and benefits of medication (composed of ROMI and ITAQ items).
	Mental state: not clinically important improvement (< 50% improvement on the PANSS), PANSS (positive, negative and general subscales).



Uzenoff 2007 (Continued)

Compliance: poor compliance (attended less than 6 sessions of therapy).

Behaviour: treatment attitude (ITAQ).

Quality of life: QLS.

Notes

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomised",' no further details reported.
Allocation concealment (selection bias)	Unclear risk	No information reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Assessments were conducted at baseline, mid treatment (3 months), and posttreatment (6 months) by interviewers blind to treatment condition."
Incomplete outcome data (attrition bias) All outcomes	Low risk	All outcome analyses were completed using a modified intent-to-treat sample (N = 19) consisting of individuals who completed both a baseline assessment and at least 1 follow-up assessment, and who had attended at least 1 session of their assigned intervention".
		"Of the 24 participants randomised to receive treatment, 3 participants did not attend and follow-up assessments".
		"Of the 24 participants randomized to receive treatment, a total of 5 participants were excluded from outcome data analyses"
Selective reporting (reporting bias)	Low risk	All expected outcomes are reported.
Other bias	Low risk	The study appears to be free of other sources of bias.

Wirshing 1991

Methods	Allocation: randomised. Blindness: none. Duration: 2 years.
	Design: 2x2 factorial.
	Setting: inpatient & outpatient.
	Country: USA.
Participants	Diagnosis: schizophrenia (DSM-III-R). N = 80. Age: mean~38 years.



Wirshing 1991 (Continued)	Sex: all male. History: stabilised for 2 months on low dose fluphenazine decanoate, no organic brain disorder, no mental retardation, no severe alcohol or substance abuse, no history of suicidal or homicidal behaviour, mean duration of illness~13 years. Exclusions: not reported.
Interventions	 Supportive group psychotherapy: dose 90 minutes twice weekly for 6 months, 90 minutes weekly for up to 18 months; goal-setting, information provision. N = 37. Behaviourally-oriented social skills training: dose 90 minutes twice weekly for 6 months, 90 minutes weekly for up to 18 months; group setting. N = 43.
Outcomes	Unable to use - Mental state: BPRS, SANS, exacerbation, severity of exacerbation, prodrome, latency from prodrome to exacerbation (data not reported). General functioning: SAS-II, degree of life disruption during an exacerbation (data not reported), symptom management skills test, medication management skills test (scale unpublished). Amount of medication needed to control an exacerbation (data not reported).
Notes	2x2 factorial design: first two conditions were behaviourally-oriented skills training program versus supportive psychotherapy control group; the other two conditions were oral neuroleptic supplementation versus placebo given in double blind fashion for the duration of each prodromal period.
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomly assigned" no further details reported.
Allocation concealment (selection bias)	Unclear risk	Details of method not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No information reported.
Blinding of outcome assessment (detection bias) All outcomes	High risk	"The rater was a clinician who was not blind to the psychosocial treatment condition."
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Losses to follow-up not reported.
Selective reporting (reporting bias)	High risk	Reports results incompletely, data not reported for BPRS, SANS, exacerbation, severity of exacerbation, prodrome, latency from prodrome to exacerbation, SAS-II, Amount of medication needed to control an exacerbation, and degree of life disruption during an exacerbation.
Other bias	Low risk	"This project was supported in part by research grants from the Veterans Administration Medical Research Service and the National Institute of Mental Health (MH141573 and MH30911)".

Diagnosis

RDC - Research Diagnostic Criteria



DSM - Diagnostic and Statistical Manual

ICD - International Classification of Diseases

Global state

GAS - Global Assessment Scale

CGI - Clinical Global Improvement

HoNOS - Health of the Nation Outcome Scales

Mental state

BDI - Beck Depression Inventory

BHS - Beck Hopelessness Scale

BPRS - Brief Psychiatric Rating Scale

BVAQ-R - Revised Beliefs About Voices Questionnaire

CESD - Center for Epidemiological Studies-Depression Scale

CPRS - Comprehensive Psychiatric Rating Scale

IMPS - Inpatient Multidimensional Psychiatric Scale

KGV - Psychiatric Assessment Scale

MADRS - Montgomery-Asberg Depression Rating Scale

NOSIE-30 - Nurses Observation Scale for Inpatient Evaluation

PANSS - Positive and Negative Syndrome Scale

PSE - Present State Examination

PSYRATS - Psychotic Symptom Rating Scale

SANS - Schedule for Assessment of Negative Symptoms

SAPS - Schedule for Assessment of Positive Symptoms

TLC - Thought, Language and Communication

Medication compliance

MCS - Medication Compliance Survey

ROMI - Rating of Medication Influences

Attitude to medication

AMQ - Attitudes to Medication Questionnaire

Insight

BCIS - Beck Cognitive Insight

ITAQ - Insight and Treatment Attitudes

SAI - Schedule for Assessment of Insight (SAI)

General functioning

AIPSS - Assessment of Interpersonal Problem-Solving Skills

GAF - Global Assessment of Function disability scale

IMPS - Inpatient Multidimensional Psychiatric Scale

KAS - Katz Adjustment Scales

LSP - Life Skills Profile

SAS II - Social Adjustment Scale II

SAS-SR - Social Adjustment Scale - Self Report

SFS - Social Functioning Scale

WAIS - Wechsler Adult Intelligence Scale

Behaviour

SBAS - Social Behaviour Adjustment Schedule

Satisfaction with treatment

CSQ - Client Service Satisfaction Questionnaire

NART - National Adult Reading Test

Quality of Life

GHQ - Global Health Questionnaire

RSES - Rosenberg Self-Esteem Scale

QLS - Quality of Life Scale

General

CBT - cognitive behavioural therapy

ST - supportive therapy

TAU - treatment as usual

Characteristics of excluded studies [ordered by study ID]



Study	Reason for exclusion	
Aberg-Wistedt 1995	Allocation: randomised. Participants: people with schizophrenia or other long-term psychotic disorders. Interventions: case management group vs standard services, not supportive therapy or supportive care.	
Abramowitz 1989	Allocation: not randomised.	
Addington 2011	Allocation: randomised. Participants: people with a high risk of developing psychosis.	
Anderson 1982	Allocation: randomised. Participants: people with schizophrenia. Interventions: family therapy vs social skills training, not supportive therapy or supportive care.	
Andres 1998	Allocation: not randomised.	
Andres 2000	Allocation: not randomised.	
Armstrong 1991	Allocation: randomised. Participants: people with major affective disorder (56%) or schizophrenia (29%) or other psychiatric disorders (16%). Interventions: life skills program vs supportive psychotherapeutic milieu. Outcomes: no data were reported separately for people with schizophrenia.	
Baker 1994	Allocation: randomised. Participants: people with schizophrenia. Interventions: problem solving training sessions vs no problem solving training, not supportive therapy or supportive care.	
Beal 1977	Allocation: randomised. Participants: people with schizophrenia. Interventions: activities group vs remotivation group vs social living group vs waiting list control group, not supportive therapy or supportive care.	
Beard 1975	Allocation: randomised. Participants: people with chronic psychiatric disorders. Interventions: group therapy sessions vs standard care, not supportive therapy or supportive care.	
Beard 1978	Allocation: randomised. Participants: people with schizophrenia (approximately 90%). Interventions: reaching out service vs community placement service vs other community facilities, not supportive therapy or supportive care.	
Becker 1998	Allocation: randomised. Participants: people with severe mental illness (44.4% schizophrenia and related psychotic disorders, 49.2% bipolar disorder and other severe mood disorders, 6.3% other diagnoses). Interventions: employment program with individual placement and support vs group skills training, not supportive therapy or supportive care.	
Beebe 2001	Allocation: randomised. Participants: people with schizophrenia. Interventions: telephone nursing intervention vs informational calls, not supportive therapy or supportive care.	
Bell 1993a	Allocation: randomised. Participants: people with schizophrenia, schizoaffective disorder.	



Study	Reason for exclusion	
	Interventions: paid work participation vs unpaid work participation, not supportive therapy or supportive care.	
Bell 1993b	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: paid work participation vs unpaid work participation, not supportive therapy or supportive care.	
Bell 1998	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions:paid work participation vs unpaid work participation, not supportive therapy or sup portive care.	
Bell 2001	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: paid work participation vs unpaid work participation, not supportive therapy or supportive care.	
Bell 2003	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: paid work participation with behavioural intervention vs paid work participation with usual support conditions, not supportive therapy or supportive care.	
Bellack 1986	Allocation: randomised. Participants: people with schizophrenia. Interventions: day hospital treatment vs day hospital treatment and social skills training, all participants received supportive care.	
Bond 1995	Allocation: randomised. Participants: people with serious mental illness (66% schizophrenia or schizoaffective disorder). Interventions: accelerated approach to supported employment vs gradual approach, not supportive therapy or supportive care.	
Brown 1983	Allocation: randomised. Participants: people with schizophrenia. Interventions: life skills vs rehabilitation programme, not supportive therapy or supportive care	
Buchkremer 1997	Allocation: randomised. Participants: people with schizophrenia. Interventions: psychoeducational medication management training vs cognitive psychotherapy velocities between the common supportive care.	
Bush 1990	Allocation: randomised. Participants: people with schizophrenia (86%) or bipolar disorder (7%) or personality disorder (7%). Interventions: intensive support from case managers vs less intensive support, not supportive therapy or supportive care.	
Canning 1997	Allocation: randomised. Participants: people with a serious psychiatric illness. Interventions: psychotherapy + family support versus psychotherapy alone.	
Carra 2010	Allocation: randomised. Participants: family members of people with schizophrenia.	
Castelein 2008	Allocation: randomised. Participants: people with psychosis. Interventions: peer support vs waiting list.	



Study	Reason for exclusion	
	Outcomes: no usable data.	
Castelein 2008a	Allocation: randomised. Participants: people with schizophrenia. Interventions: peer to peer support groups vs waiting list.	
Castilla-Puentes 2002	Allocation: randomised. Participants: people with schizophrenia. Interventions: supportive talking/counselling versus group music therapy. Outcomes: no usable data.	
Chien 2008	Allocation: randomised. Participants: family caregivers of people with schizophrenia. Interventions: mutual support group vs standard treatment.	
Chinman 2010	Allocation: not randomised.	
Clark 1998	Allocation: randomised. Participants: people with severe mental illness (46.9% schizophrenia or related psychotic disorders, 42.7% bipolar disorder or other severe mood disorders, 10.5% other disorders). Interventions: individual placement and support for employment vs group skills training, not supportive therapy or supportive care.	
Cook 2011	Allocation: randomised. Participants: persons with serious mental illness, < 50% schizophrenia. Interventions: Building Recovery of Individual Dreams and Goals through Education and Support (BRIDGES) vs waiting list.	
Cook 2012	Allocation: randomised.	
	Participants: persons with serious mental illness, < 50% schizophrenia.	
	Interventions: Building Recovery of Individual Dreams and Goals through Education and Support (BRIDGES) vs waiting list.	
Curtis 1992	Allocation: randomised. Participants: people who were being discharged from psychiatric hospital (44.7% schizophrenia). Interventions: multidisciplinary intensive outreach case management vs less intensive community support system case management services vs routine aftercare with no case management, not supportive therapy or supportive care.	
Curtis 1996	Allocation: randomised. Participants: people who had received inpatient psychiatric care (38.4% schizophrenia). Interventions: intensive outreach case management vs standard aftercare services, not supportive therapy or supportive care.	
Czobor 1995	Allocation: randomised. Participants: people with schizophrenia. Interventions: risperidone vs placebo, not supportive therapy or supportive care.	
D'Ercole 1997	Allocation: randomised. Participants: people with major psychiatric disorder (38% schizophrenia). Interventions: intensive outreach case management vs standard aftercare, not supportive therapy or supportive care.	
Daumit 2010	Allocation: randomised. Participants: persons with severe mental illness, 25% schizophrenia. Interventions: group exercise vs group exercise plus peer support.	



Study	Reason for exclusion
Dixon 2002	Allocation: randomised. Participants: people with severe mental illness. Interventions: individual placement and support vs enhanced vocational rehabilitation, not supportive therapy or supportive care.
Donlon 1973	Allocation: randomised. Participants: people with schizophrenia. Interventions: group socialisation, refreshments and support versus individual supportive psychotherapy; both interventions are supportive therapy or supportive care.
Drake 1993	Allocation: randomised. Participants: people with severe mental illness and substance use disorder. Interventions: social network treatment vs CBT, not supportive therapy or supportive care.
Drake 1996	Allocation: randomised. Participants: people with severe mental illness (46.9% schizophrenia and related psychotic disorders, 42.7% bipolar disorder and other severe mood disorders, 10.5% other disorders). Interventions: group skills training vs individual placement and support, not supportive therapy or supportive care.
Drake 1999	Allocation: randomised. Participants: people with severe mental disorders (67% schizophrenia). Interventions: individual placement and support vs enhanced vocational rehabilitation, not supportive therapy or supportive care.
Drury 1996a	Allocation: randomised. Participants: people with non-affective psychosis. Interventions: individual and group cognitive therapy vs matched hours of therapy, not supportive therapy or supportive care.
Drury 1996b	Allocation: randomised. Participants: people with non-affective psychosis. Interventions: individual and group cognitive therapy vs matched hours of therapy, not supportive therapy or supportive care.
Drury 2000	Allocation: randomised. Participants: people with non-affective psychosis. Interventions: cognitive therapy programme vs recreational activities and support, not supportive therapy or supportive care.
Eack 2007	Allocation:randomised. Participants: people with schizophrenia. Interventions: Enriched Supportive therapy (psycho-education and training) vs CBT.
Eack 2010	Allocation:randomised. Participants: people with schizophrenia. Interventions: Enriched Supportive therapy (psycho-education and training) vs CBT.
Eack 2011	Allocation:randomised. Participants: people with schizophrenia. Interventions: Enriched Supportive therapy (psycho-education and training) vs CBT.
Eack 2012	Allocation: randomised. Participants: people with schizophrenia. Interventions: Enriched Supportive therapy (psycho-education and training) vs CBT.
Essock 1995	Allocation: randomised.



Study	Reason for exclusion		
	Participants: people with serious mental disorder (67% schizophrenia or schizoaffective disorder). Interventions: assertive community treatment vs case management, not supportive therapy or supportive care.		
Field 1997	Allocation: not randomised.		
Ford 1995	Allocation: randomised. Participants: people with mental illness (82% schizophrenia). Interventions: case management vs standard psychiatric services, not supportive therapy or supportive care.		
Forsyth 1961	Allocation: not randomised.		
Franklin 1987	Allocation: randomised. Participants: people with a minimum of two discharges from a mental hospital (56% schizophrenia). Interventions: case management services vs any services except case management, not supportive therapy or supportive care.		
Freeman 1998	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or delusional disorder. Interventions: standard care vs standard care and CBT, not supportive therapy or supportive care.		
Gaither 2003	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: cognitive behavioural intervention vs interpersonal attention control, not supportive therapy or supportive care.		
Garety 1994	Allocation: not randomised.		
Glick 1974	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.		
Glick 1975	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.		
Glick 1976a	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.		
Glick 1976b	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long		
	-term hospitalisation, not supportive therapy or supportive care.		
Glick 1976c	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.		
Glick 1977	Allocation: randomised. Participants: people with schizophrenia.		



Study	Reason for exclusion
	Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.
Glick 1979	Allocation: randomised. Participants: people admitted to a psychiatric inpatient ward (60% schizophrenia). Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.
Glynn 2001	Allocation: randomised. Participants: people with schizophrenia. Interventions: clinic skills training vs clinic skills training with manualised weekly community instruction to generalise skills to the natural environment, not supportive therapy or supportive care
Grassi 2001	Allocation: not randomised.
Grawe 1998	Allocation: randomised. Participants: people with schizophrenia. Interventions: optimal multimodal treatment vs treatment as usual, not supportive therapy or supportive care.
Haddock 2000a	Allocation: randomised. Participants: people with schizophrenia and substance misuse. Interventions: individual CBT vs family support, not supportive therapy or supportive care.
Haddock 2000b	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or delusional disorder and alcohol or drug abuse. Interventions: individual CBT vs family support, not supportive therapy or supportive care.
Hafner 1983	Allocation: randomised. Participants: people with severe and persisting psychiatric disorders, the majority did not have schizophrenia.
	Interventions: couples therapy vs individual therapy.
Haldun 2002	Allocation: randomised. Participants: people with schizophrenia. Interventions: optimal clinical management vs routine case management, not supportive therapy or supportive care.
Hannes 1974	Allocation: randomised. Participants: people with schizophrenia. Interventions: recreational therapy vs relaxation period, not supportive therapy or supportive care.
Hargreaves 1977	Allocation: randomised. Participants: people with schizophrenia. Interventions: short term hospitalisation vs long-term hospitalisation, not supportive therapy or supportive care.
Harrison-Read 2002	Allocation: randomised. Participants: people who are heavy users of inpatient psychiatric services (majority are people with schizophrenia or schizotypal disorders or delusional disorders). Interventions: enhanced community management vs standard care, not supportive therapy or supportive care.
Harvey 2002	Allocation: randomised. Participants: relatives of people with psychosis.



Study	Reason for exclusion
	Interventions: standard case management vs intensive case management.
Herz 1974	Allocation: randomised. Participants: people with psychiatric illness (66% schizophrenia). Interventions: individual supportive psychotherapy versus group supportive psychotherapy.
Herz 2000	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: program for relapse prevention vs treatment as usual, supportive therapy received by all participants.
Hogarty 1973	Allocation: randomised. Participants: people with schizophrenia. Interventions: major role therapy and chlorpromazine vs major role therapy and placebo vs chlorpromazine vs placebo, not supportive therapy or supportive care.
Hogarty 1977	Allocation: randomised. Participants: people with schizophrenia. Interventions: major role therapy and chlorpromazine vs major role therapy and placebo vs chlorpromazine vs placebo, not supportive therapy or supportive care.
Hogarty 1984	Allocation: randomised. Participants: people with schizophrenia. Interventions: major role therapy and chlorpromazine vs major role therapy and placebo vs chlorpromazine vs placebo, not supportive therapy or supportive care.
Hogarty 1986a	Allocation: randomised. Participants: people with schizophrenia. Interventions: family treatment and medication vs social skills training and medication vs family treatment + social skills training + medication vs medication.
Hogarty 1986b	Allocation: randomised. Participants: people with schizophrenia. Interventions: family treatment and medication vs social skills training and medication vs family treatment + social skills training + medication vs medication.
Hogarty 1986c	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: family treatment and medication vs social skills training and medication vs family treatment + social skills training + medication vs medication, not supportive therapy or supportive care.
Hogarty 1988	Allocation: randomised. Participants: people with schizophrenia. Interventions: standard dose of fluphenazine decanoate vs minimal dose, not supportive therapy or supportive care.
Hornung 1998	Allocation: randomised. Participants: people with schizophrenia. Interventions: psychoeducational medication management vs cognitive psychotherapy vs key person counselling, not supportive therapy or supportive care.
Hornung 1999	Allocation: randomised. Participants: people with schizophrenia. Interventions: psychoeducational medication management vs cognitive psychotherapy vs key person counselling, not supportive therapy or supportive care.
Hurlburt 1996	Allocation: randomised.



Study	Reason for exclusion
	Participants: people with severe mental illness (55.4% schizophrenia). Interventions: access to rent subsidy certificate vs no access to rent subsidy certificate vs tradition al services vs comprehensive services, not supportive therapy or supportive care.
Huxley 2001	Allocation: randomised. Participants: people with psychosis. Interventions: intensive case management vs standard case management, not supportive therapy or supportive care.
Isrctn50487713 2010	Allocation: partially randomised (participants able to choose their preferred treatment option or elect to be randomised to a treatment option. Participants: people with schizophrenia. Interventions: peer support and CBT vs peer support, CBT and group sessions.
Isrctn69299093 2011	Allocation: randomised. Participants: family members of people with psychotic symptoms. Interventions: Relatives' Education and Coping Toolkit (REACT) vs CBT oriented self management intervention.
Isrctn96754763 2011	Allocation:randomised. Participants: staff working on acute mental health wards. Interventions: modified cognitive analytic therapy approach to clinical supervision vs inactive control arm.
Issakidis 1999	Allocation: randomised. Participants: people with schizophrenia or bipolar disorder. Interventions: intensive case management vs standard case management, not supportive therapy or supportive care.
Jones 1994	Allocation: randomised. Participants: people with a DSM-III-R Axis I diagnosis (63% schizophrenia). Interventions: critical time intervention vs usual treatment services for homeless people with severe mental illness, not supportive therapy or supportive care.
Kaplan 2011	Allocation:randomised. Participants: people with schizophrenia. Interventions: experimental peer support listserv (group distribution email list) vs experimental peer support online bulletin vs waiting list control group.
Karon 1969	Allocation: randomised. Participants: people with schizophrenia. Interventions: supportive therapy + medication versus 'active' psychoanalytic therapy + no medication versus 'ego-analytic' psychoanalytic therapy + medication as an adjunct; intervention groups received different medication regimes therefore results are confounded.
Kaufmann 1995	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or major affective disorder. Interventions: self help employment centre services vs customary community service, not supportive therapy or supportive care.
Keith 1977	Allocation: randomised. Participants: rehabilitation service clients. Interventions: innovative training program vs usual service + outcome measures vs usual services only, not supportive therapy or supportive care.
Kern 2002	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder.



Study	Reason for exclusion
	Interventions: work training by errorless learning vs conventional instruction, not supportive therapy or supportive care.
Keshavan 2011	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder; criteria not reported.
	Interventions: cognitive enhancement therapy vs. supportive therapy.
	Outcomes: no usable data, abstract only.
Kim 1997	Allocation: not randomised.
Klingberg 1999	Allocation: randomised. Participants: people with schizophrenia. Interventions: psychoeducational medication management training vs cognitive psychotherapy vs key-person counselling, not supportive therapy or supportive care.
Klingberg 2010	Allocation: randomised. Participants: schizophrenia, schizophreniform disorder, schizoaffective disorder, delusional disorder.
	Interventions: CBT vs non-specific supportive therapy.
	Outcomes: no usable data.
Koivunen 2010	Allocation:randomised. Participants: nurses on acute psychiatric wards. Interventions: web-based patient support system vs standard patient education sessions with written material.
Kris 1965	Allocation: randomised. Participants: people who have previously been in a psychiatric hospital, with severe psychotic relapse. Interventions: day hospital vs state mental hospital, not supportive therapy or supportive care.
Kuipers 1998	Allocation: randomised. Participants: people with schizophrenia or delusional disorder or schizoaffective disorder. Interventions: CBT vs standard care, not supportive therapy or supportive care.
Lecomte 1999	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: self-esteem module vs standard care, not supportive therapy or supportive care.
Lehman 1993	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or major affective disorder and substance use disorder. Interventions: innovative group and intensive case management + usual community mental health centre and rehabilitation services vs usual community mental health centre and rehabilitation services, not supportive therapy or supportive care.
Lehman 2001	Allocation: randomised. Participants: people with severe mental illness (70% schizophrenia). Interventions: Individual Placement and Support (IPS) employment program vs comparison psychosocial program, not supportive therapy or supportive care.
Lehman 2002a	Allocation: randomised. Participants: people with severe mental illness (75% chronic psychoses).



Study	Reason for exclusion
	Interventions: Individual Placement and Support (IPS) employment program vs psychosocial rehabilitation program, not supportive therapy or supportive care.
Lehman 2002b	Allocation: randomised. Participants: people with severe mental illness (75% chronic psychoses). Interventions: Individual Placement and Support (IPS) employment program vs psychosocial rehabilitation program, not supportive therapy or supportive care.
Levene 1989	Allocation: randomised. Participants: people with schizophrenia. Interventions: focal family therapy vs supportive management counselling, not supportive therapy or supportive care.
Lewis 2000	Allocation: randomised. Participants: people with schizophrenia and substance use. Interventions: family support vs CBT, not supportive therapy or supportive care.
Lewis 2001a	Allocation: randomised. Participants: people with schizophrenia. Interventions: CBT + drug treatment vs drug treatment, not supportive therapy or supportive care.
Lindenmayer 1995	Allocation: randomised. Participants: people with schizophrenia. Interventions: placebo vs haloperidol vs risperidone, not supportive therapy or supportive care.
Linszen 1994	Allocation: randomised. Participants: people with schizophrenia or schizophreniform disorder or schizoaffective disorder or other schizophrenia-like psychotic disorder. Interventions: individual treatment versus family treatment + individual treatment.
Linszen 1996	Allocation: randomised. Participants: people with schizophrenia. Interventions: individual orientated intervention vs individual and family orientated intervention, not supportive therapy or supportive care.
Linszen 1997	Allocation: randomised. Participants: people with schizophrenia or schizophreniform disorder or schizoaffective disorder or other psychotic disorder. Interventions: individual treatment versus family treatment + individual treatment, not supportive therapy or supportive care.
Linszen 1998	Allocation:randomised. Participants: people with schizophrenia. Interventions: individual treatment versus family treatment + individual treatment, not supportive therapy or supportive care.
Linszen 2009	Allocation:randomised. Participants: people with schizophrenia and parents of schizophrenic patients. Interventions: Continuity of Treatment by professionals Specialized in the treatment of early schizophrenia (CST) vs Continuity of Specialized Treatment plus Parent groups (CSTpP) vs Continuity of Treatment as Usual (CTU).
Macias 1994	Allocation: randomised. Participants: people with serious mental illness (46% schizophrenia, 22% affective disorder). Interventions: team case management vs rehabilitation program, not supportive therapy or supportive care.
Marder 1994a	Allocation: randomised.



Study	Reason for exclusion
	Participants: people with schizophrenia. Interventions: risperidon vs placebo, not supportive therapy or supportive care.
Marder 1994b	Allocation: randomised. Participants: people with schizophrenia. Interventions: oral fluphenazine vs placebo, not supportive therapy or supportive care.
Marder 2001b	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: Individual Placement and Support + Psychoeducational Skills Training vs Individual Placement and Support, not supportive therapy or supportive care.
Marshall 1985	Allocation: randomised. Participants: family members of people with schizophrenic disorder.
	Interventions: skills training and education vs education only.
May 1976a	Allocation: randomised. Participants: people with schizophrenia. Interventions: individual psychotherapy vs ataraxic drugs vs individual psychotherapy + ataraxic drugs vs ECT vs milieu treatment alone, not supportive therapy or supportive care.
McEvoy 1994	Allocation: randomised. Participants: people with schizophrenia. Interventions: risperidone vs haloperidol vs placebo, not supportive therapy or supportive care.
McFarlane 1992	Allocation: not randomised.
McFarlane 2002	Allocation: randomised. Participants: people with disabling psychiatric disorder. Interventions: Mental Health Employers Consortium vs conventional methods of supported employment, not supportive therapy or supportive care.
McGorry 1999	Allocation: randomised. Participants: people with an at risk mental state.
	Interventions: supportive psychotherapy alone (control group) or specific cognitive psychotherapy plus low dose risperidone. Outcome: no usable data, abstract only.
McGorry 2000	Allocation: randomised. Participants: people with schizophrenia.
	Interventions: risperidone and cognitive therapy vs supportive case management. Outcome: no usable data, abstract only.
McInnis 1967	Allocation: randomised. Participants: people with schizophrenia. Interventions: positive extrinsic reinforcement vs negative extrinsic reinforcement, not supportive therapy or supportive care.
Meisler 2002	Allocation: not stated. Participants: people with severe mental illness (50% schizophrenia related disorders). Interventions: Program of Assertive Community Treatment-PACT vs Individual Placement and Support-IPS (ACT-IPS), not supportive therapy or supportive care.
Meister 2010	Allocation: randomised. Participants: psychosis and substance use disorder; criteria not reported.



Study	Reason for exclusion
	Interventions: group-based motivational behavioural therapy for drug abuse vs supportive treatment for addiction recovery.
	Outcomes: no usable data.
Miknyak 2001	Allocation: not stated. Participants: people with schizophrenia. Interventions: risperidone vs fluanxol-depot vs risperidone + fluanxol-depot, not supportive therapy or supportive care.
Morse 1997	Allocation: randomised. Participants: people with a serious DSM-III-R axis I diagnosis (66% schizophrenia). Interventions: broker care management vs assertive community treatment vs assertive community treatment + support from community workers, not supportive therapy or supportive care.
Mosher 1975	Allocation: not randomised.
Mueser 2001a	Allocation: randomised. Participants: people with schizophrenia or other severe mental illness. Interventions: Individual Placement and Support (IPS) vs psychiatric rehabilitation program vs standard services, not supportive therapy or supportive care.
Mueser 2001b	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or schizophreniform disorder. Interventions: supportive family treatment vs applied family therapy, not supportive therapy or supportive care.
NCT 2005	Allocation: randomised. Participants: people with schizophrenia. Interventions: Health Care Management and Supported Rehabilitation (includes skills training) vs standard treatment.
NCT 2005a	Allocation: randomised. Participants: people with schizophrenia and their families. Interventions: online advice from healthcare professionals, social support, and information vs standard treatment.
NCT 2009	Allocation: randomised. Participants: people with schizophrenia. Interventions: support group vs mindfulness meditation.
NCT 2009a	Allocation: randomised. Participants: family of people with schizophrenia. Interventions: family led mutual support group vs psycho-education and standard treatment.
NCT 2012a	Allocation: randomised. Participants: people with schizophrenia. Interventions: Enriched Supportive therapy (psycho-education and training) vs Cognitive Enhance-ment therapy.
Nelson 2007	Allocation: randomised. Participants: ultra high risk criteria for psychotic disorder. Interventions: low- dose risperidone + intensive CBT-based psychological treatment vs placebo + intensive CBT-based psychological treatment vs placebo + control psychological treatment (supportive therapy). Outcomes: no usable data, abstract only.
Nugter 1997a	Allocation: randomised.



Study	Reason for exclusion
	Participants: people with schizophrenia or schizophreniform disorder or schizoaffective disorder or other psychotic disorders. Interventions: individual treatment versus individual treatment + behavioural family treatment.
Nugter 1997b	Allocation: randomised. Participants: people with schizophrenia or schizophreniform disorder or schizoaffective disorder or other psychotic disorders. Interventions: individual treatment vs individual treatment + family treatment, not supportive therapy or supportive care.
O'Donnell 2002	Allocation: randomised. Participants: people with schizophrenia. Interventions: compliance therapy vs non-specific counselling therapy, not supportive therapy or supportive care.
Ogrodniczuk 2000	Allocation: randomised. Participants: people with major depression or dysthymia or personality disorder.
Ohlenschlaeger 2002	Allocation: randomised. Participants: people with schizophrenia or schizotypal disorders or delusional disorders. Interventions: standard treatment vs hospital based rehabilitation, not supportive therapy or supportive care.
Philipps 2001	Allocation: randomised. Participants: young people identified as being at high risk of psychosis. Interventions: combined medical and psychological (specific) approach vs supportive (non-specific) case management.
Phillips 1999	Allocation: randomised. Participants: young people identified as being at high risk of psychosis. Interventions: combined medical and psychological approach vs supportive case management.
Pickett 1997	Allocation: not randomised.
Pushkaryova 1999	Allocation: randomised. Participants: people with psychosis. Interventions: enterosorption of silicorganic enterosorbent vs placebo, not supportive therapy or supportive care.
Razali 1995	Allocation: randomised. Participants: people with schizophrenia. Interventions: counselled by trained hospital pharmacist vs no counselling, not supportive therapy or supportive care.
Rector 2002	Allocation: randomised. Participants: people with schizophrenia. Interventions: cognitive therapy vs standard care, not supportive therapy or supportive care.
Reynolds 2002	Allocation: randomised. Participants: people discharged from a psychiatric inpatient unit. Interventions: transitional discharge model vs standard discharge care, not supportive therapy or supportive care.
Ritch 2001	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder.



Study	Reason for exclusion
	Interventions: Cognitive Adaptation Training vs a condition designed to control for therapist time and provide environmental changes not associated with cognitive deficits vs assessment only., not supportive therapy or supportive care.
Ritch 2002	Allocation: randomised. Participants: people with high service utilisation. Interventions: Assertive Community Treatment vs Cognitive Adaptation Training, not supportive therapy or supportive care.
Ro-Trock 1977	Allocation: randomised. Participants: adolescents with schizophrenic reaction (57%), adolescent adjustment reaction or drug problem Interventions: family therapy vs individual therapy, not supportive therapy or supportive care.
Rosenheck 2003	Allocation: randomised. Participants: people with major psychiatric disorder or an alcohol or drug abuse disorder or both (~10% serious psychiatric disorders, ~50% alcohol or drug disorders, ~35% dual diagnoses, ~5% other psychiatric diagnoses).
	Interventions:supported housing program + intensive case management vs case management only vs standard care.
Rosenthal 2000	Allocation: randomised. Participants: people with schizophrenia and substance abuse. Interventions: outpatient group therapy vs group therapy plus Targeted Assertive Outreach visits, all participants received group therapy.
Ruan 2008	Allocation:randomised. Participants: people with schizophrenia. Interventions: clozapine vs clozapine plus fluoxetine, both groups received supportive therapy.
Serok 1984	Allocation: randomised. Participants: people with schizophrenia. Interventions: Gestalt therapy vs care as usual, not supportive therapy or supportive care.
Shi 2000	Allocation: not clear. Participants: people with schizophrenia. Interventions: family therapy vs control, not supportive therapy or supportive care.
Shin 2002	Allocation: randomised. Participants: people with schizophrenia. Interventions: psycho-educational group programme + individual supportive therapy vs individual supportive therapy alone.
Slavinsky 1982	Allocation: randomised (but allocation concealment inadequate).
Smith 1999	Allocation: randomised. Participants: people with schizophrenia. Interventions: supportive group psychotherapy vs skills training. Outcomes: no usable data.
Solomon 1994a	Allocation: randomised. Participants: people with serious mental illness (85% schizophrenia). Interventions: intensive case management by an assertive community team vs intensive case management by individual case managers vs community mental health centre, not supportive therapy or supportive care.
Solomon 1994b	Allocation: randomised.



Study	Reason for exclusion	
	Participants: people with serious mental illness (85.9% schizophrenia). Interventions: consumer case managers vs non-consumer case management, not supportive therapy or supportive care.	
Solomon 1995a	Allocation: randomised. Participants: people with serious mental illness (86% schizophrenia). Interventions: consumer case managers vs non-consumer case management, not supportive therapy or supportive care, not supportive therapy or supportive care.	
Solomon 1995b	Allocation: randomised. Participants: people with schizophrenia (85%) or major affective disorder (14%) or unspecified psychotic disorder (1%). Interventions: consumer case managers vs non-consumer case management, not supportive therapy or supportive care.	
Solomon 1996	Allocation: randomised. Participants: people with serious mental illness. Interventions: consumer case managers vs non-consumer case management, not supportive therapy or supportive care.	
Stant 2011	Allocation:randomised. Participants: people with schizophrenia. Interventions: peer support group vs standard treatment.	
Stroup 2003	Allocation: randomised. Participants: people with schizophrenia. Interventions: typical antipsychotics vs atypical antipsychotics, not supportive therapy or supportive care.	
Sun ShuMin 2007	Allocation: randomised. Participants: people with schizophrenia. Interventions: social support therapy (includes psycho-education, delivered by a group of people) vs standard treatment.	
Tarrier 2000a	Allocation: randomised. Participants: people with schizophrenia and substance misuse. Interventions: family support vs cognitive service, not supportive therapy or supportive care.	
Tarrier 2000d	Allocation: randomised. Participants: support workers with care load of people with schizophrenia. Interventions: training in family support vs not reported (trial register).	
Telles 1995	Allocation: randomised. Participants: people with schizophrenia. Interventions: standard case management vs behavioural family management. Outcomes: no usable data.	
Test 1991	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder or schizotypal personality. Interventions: training in community living vs case management, not supportive therapy or supportive care.	
Theilemann 1993	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: "place-train" supported employment strategies vs employment strategies along with intensive interventions, not supportive therapy or supportive care.	



Study	Reason for exclusion	
Tong Shouming 2010	Allocation: not randomised.	
Toprac 2002	Allocation: randomised. Participants: people with serious mental illness. Interventions: integrated supported employment vs group skills training, not supportive therapy or supportive care.	
Torrey 2000	Allocation: randomised. Participants: people with severe mental disorders (46.9% chronic psychotic illnesses, 42.7% severe affective disorders, 10.4% other disorders). Interventions: XX vs YY, not supportive therapy or supportive care.	
Tsang 2001	Allocation: randomised. Participants: people with schizophrenia. Interventions: social and job skills training with support vs social and job skills training without support vs assessment only, not supportive therapy or supportive care.	
Tyrer 1995	Allocation: randomised. Participants: people on a register for vulnerable psychiatric patients (54% schizophrenia). Interventions: supervision by nominated key workers vs standard care, not supportive therapy or supportive care.	
Vaughan 1992	Allocation: randomised. Participants: people with schizophrenia. Interventions: relatives' counselling + standard after-care versus standard after-care alone; not supportive therapy or supportive care.	
Velligan 1999	Allocation: randomised. Participants: people with schizophrenia or schizoaffective disorder. Interventions: cognitive adaptation training vs attention control vs follow-up only, not supportive therapy or supportive care.	
Walker 1969	Allocation: randomised. Participants: psychiatric inpatients (50% schizophrenic reaction). Interventions: community hospital industrial rehabilitation placement vs waiting list, not supportive therapy or supportive care.	
Wallace 1985	Allocation: randomised. Participants: people with schizophrenia. Interventions: intensive social skills training vs holistic health therapy, not supportive therapy or supportive care.	
Wallace 2001	Allocation: randomised. Participants: people with a serious and persistent mental illness. Interventions: Individual Placement and Support vs Individual Placement and Support + workplace fundamentals module, not supportive therapy or supportive care.	
Weinman 1974	Allocation: randomised. Participants: people with schizophrenia or functional psychosis. Interventions: community placement vs hospital based socio-environmental condition, not supportive therapy or supportive care.	
Wojciechowska 2001	Allocation: not randomised.	
Wolkon 1971	Allocation: randomised. Participants: people recently discharged from psychiatric hospital with schizophrenia (78%) or psychotic disorder or non-psychotic disorder.	



Study	Reason for exclusion	
	Interventions: social rehabilitation program vs control group, not supportive therapy or supportive care.	
Wuerker 2002	Allocation: randomised. Participants: people with schizophrenia and their relatives. Interventions: supportive family case management vs applied family management, not supportive therapy or supportive care.	
Yung 2008	Allocation: randomised. Participants: people with ultra high risk of psychotic disorder. Interventions: low-dose risperidone plus CBT vs placebo plus CBT vs placebo plus supportive therapy.	
Yung 2011	Allocation: randomised. Participants: people with ultra high risk of psychotic disorder. Interventions: cognitive therapy plus risperidone vs cognitive therapy plus placebo vs supportive therapy plus placebo.	
Ziedonis 1997	Allocation: not randomised.	

CBT - cognitive behavioural therapy

DSM-III - Diagnostic and Statistical Manual third edition

 ${\sf ECT-electroconvulsive\ the rapy}$

Characteristics of studies awaiting assessment [ordered by study ID]

Bechdolf 2012

Methods	Allocation: randomised. Blindness: unclear. Duration: 6 months.
	Design: parallel.
	Setting: inpatient.
	Country: Germany.
Participants	Diagnosis: schizophrenia and substance abuse disorder. N = 60. Age: unclear. Sex: unclear. History: unclear. Exclusions: unclear.
Interventions	1. Motivational interviewing: four sessions. N = unclear.
	2. Supportive therapy: four sessions. N = unclear.
Outcomes	Unclear.
Notes	Paper in German, awaiting translation



Blankertz 1997	
Methods	Unclear.
Participants	Unclear.
Interventions	Unclear.
Outcomes	Unclear.
Notes	Waiting for a translation of the paper

Falloon 1983

Methods	Unclear.
Participants	Unclear.
Interventions	Unclear.
Outcomes	Unclear.
Notes	No full text available.

Characteristics of ongoing studies [ordered by study ID]

ACTRN12610000241033 2010

Trial name or title	A pilot project investigating the feasibility and effects of a 6-month peer support program commencing 3 months prior to discharge from a specialist first-episode psychosis treatment centre.	
Methods	Randomised controlled trial (computer-generated random numbers). Parallel assignment. Outcome assessors: blinded.	
Participants	First-episode psychosis.	
Interventions	Intervention: Individual peer support. Control: Standard care.	
Outcomes	Primary outcome: Increased service satisfaction, as measured by the Verona Service Satisfaction Scale. Secondary outcome: Reduced risk of suicide, as measured by The Reynolds Suicidal Ideation Questionnaire.	
Starting date	1/6/2010.	
Contact information	Jo Robinson Orygen Youth Health Research Centre 35 Poplar Road Parkville, Vic, 3052 Australia +61 3 9342 2866 +61 3 9342 2941 jo.robinson@mh.org.au	



ACTRN12610000241033 2010 (Continued)

Notes

ACTRN12612000974808 2012

Trial name or title	Peer-delivered support intervention for people who hear voices: Pilot randomised controlled trial	
Methods	Randomised controlled trial using computerised sequence generation.	
	Blinded by masking.	
	Crossover assignment.	
Participants	18-65 years old, Auditory verbal hallucinations at least weekly over the past six months associated with significant distress or disability.	
Interventions	Intervention: 12 weekly one hour one-to-one sessions of a support intervention for hearing voices (auditory verbal hallucinations) from a peer mental health worker who has had personal lived experience of hearing voices themselves. The intervention group will receive the intervention alongside treatment as usual (TAU) in the 3-month period immediately following randomisation.	
	Control: Treatment as usual (wait list). The control group will receive the intervention after a 3-month treatment as usual wait list period. Treatment as usual will involve the person's usual men tal health care which will ordinarily include prescription of antipsychotic medication, plus meetings with mental health workers and possible attendance at rehabilitative and recovery-oriented mental health service programmes.	
Outcomes	Primary outcome: Subjective Experiences of Psychosis Scale.	
	Secondary outcome: Recovery Assessment Scale.	
Starting date	1/11/2012	
Contact information	Ms Indigo Daya, Voices Vic 15 Cromwell Road South Yarra VIC 314, Australia,	
	+61 3 9692 9562,	

Bennett 2003

Trial name or title	Treating substance abuse in schizophrenia: Preliminary analysis of data from a randomised treatment trial.
Methods	
Participants	People with schizophrenia.
Interventions	Behavioural treatment for substance abuse. supportive therapy.
Outcomes	Treatment retention, treatment utilisation, urine data.



Ctautina adata	
Starting date	
Contact information	
Notes	
Ourham 2000a	
Trial name or title	Cognitive behavioural therapy with medication resistant psychotic symptoms: a clinical trial with psychiatric nurses as therapists.
Methods	
Participants	People with schizophrenia.
Interventions	CBT. Supportive counselling.
Outcomes	To assess the efficacy of delivery of care by clinical nurse specialist.
Starting date	
Contact information	
Notes	
logarty 2001	Environmental personal treatment of sphirophropia
logarty 2001 Trial name or title	Environmental-personal treatment of schizophrenia.
logarty 2001	Environmental-personal treatment of schizophrenia.
logarty 2001 Trial name or title	Environmental-personal treatment of schizophrenia. People with schizophrenia.
logarty 2001 Trial name or title Methods	
Iogarty 2001 Trial name or title Methods Participants	People with schizophrenia. 1. Cognitive Enhancement Therapy.
Hogarty 2001 Trial name or title Methods Participants Interventions	People with schizophrenia. 1. Cognitive Enhancement Therapy. 2. Enriched Supportive Therapy. Interpersonal and vocational effectiveness, self-esteem, neuropsychological competence and
Iogarty 2001 Trial name or title Methods Participants Interventions Outcomes	People with schizophrenia. 1. Cognitive Enhancement Therapy. 2. Enriched Supportive Therapy. Interpersonal and vocational effectiveness, self-esteem, neuropsychological competence and
Hogarty 2001 Trial name or title Methods Participants Interventions Outcomes Starting date	People with schizophrenia. 1. Cognitive Enhancement Therapy. 2. Enriched Supportive Therapy. Interpersonal and vocational effectiveness, self-esteem, neuropsychological competence and
Hogarty 2001 Trial name or title Methods Participants Interventions Outcomes Starting date Contact information	People with schizophrenia. 1. Cognitive Enhancement Therapy. 2. Enriched Supportive Therapy. Interpersonal and vocational effectiveness, self-esteem, neuropsychological competence and



ISRCTN02457313	2006 (Continued)
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Methods	Randomised controlled trial. Parallel group.
Participants	Patients diagnosed with schizophrenia, psychosis.
Interventions	Intervention: Peer support group. Control: Waiting list control group.
Outcomes	Primary outcome: Quality of Life: WHO Qol Bref. Secondary outcomes: 1. Social support 2. Social network 3. Self-efficacy 4. Self esteem 5. Psychopathology.
Starting date	Feb 2003.
Contact information	Not reported.
Notes	Trial ended.

ISRCTN50487713 2011

Trial name or title	An evaluation of different levels of support in using a recovery guide for people with psychosis and the impact of choice on outcomes
Methods	Partially randomised patient preference trial. Participants may choose their preferred treatment option: Treatment As Usual, low support/high support or elect to be randomised to a treatment op tion.
Participants	Meeting ICD 10 criteria for non affective psychosis (schizophrenia,schizophreniform disorder, schizoaffective disorder, delusional disorder).
Interventions	Interventions: 1. Low support 2. High support Control: Treatment as usual.
Outcomes	Psychotic symptoms, affect, well-being and functioning.
Starting date	Not reported.
Contact information	Prof Gillian Haddock University of Manchester School of Psychological Sciences Zochonis Building Oxford Road Manchester Greater Manchester M13 9PL UNITED KINGDOM Tel: 0161 275 8756 gillian.haddock@manchester.ac.uk
Notes	The trial is reported as "closed - in follow-up".



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Trial name or title	A randomised controlled trial of individual therapy for first episode psychosis (PSTEP).
Methods	Randomised, single-blind (outcomes assessor), placebo-control, single-group assignment.
Participants	Schizophrenia, schizophreniform disorder, brief psychotic disorder, delusional disorder, schizoaffective disorder, substance induced psychotic disorder, or psychotic disorder NOS.
Interventions	CBT; befriending.
Outcomes	Social functioning; positive & negative symptoms; individual dimensions of psychotic symptoms; depression; substance use; alcohol and drug use; medication adherence adaptation to illness; self esteem; coping skills.
Starting date	September 2007
Contact information	Diane Kirsopp, BA 416-535-8501 ext 6288 diane_kirsopp@camh.net
Notes	

NCT 2007a

Trial name or title	Peer support for increasing physical activity in people with serious mental illnesses.
Methods	Randomised, single-blind (outcomes assessor), active control, parallel assignment
Participants	Patient at the Johns Hopkins Community Psychiatry program: serious mental illnesses.
Interventions	Physical activity intervention with peer support (PA + PS) Physical activity intervention without peer support (PA)
Outcomes	Primary: Cardiorespiratory fitness Secondary: Weight; waist circumference; physical activity; health status; Center for Epidemiology Depression Scale; exercise-related self-efficacy; general perceived efficacy; participation.
Starting date	March 2007.
Contact information	Gail L. Daumit, MD, MHS 410-614-6460 gdaumit@jhmi.edu
Notes	

NCT 2007b

Trial name or title	Recovery Group Pilot.
Methods	Randomised,open label, uncontrolled, parallel assignment, delayed control group and 6- and 12-month post-intervention follow-up.



NCT 2007b (Continued)	
Participants	Bipolar disorder, schizophrenia, schizoaffective disorder.
Interventions	Recovery group workbook & group sessions
Outcomes	Improvements in short-term recovery and functioning, participant attendance and satisfaction with the group sessions.
Starting date	July 2005.
Contact information	Not reported.
Notes	

NCT 2008

Trial name or title	A randomised controlled trial of individual therapy for first episode psychosis
Methods	Randomised, single-blind, active control, parallel assignment.
Participants	DSM-IV criteria for: schizophrenia, schizophreniform disorder, brief psychotic disorder, delusional disorder, schizoaffective disorder, substance induced psychotic disorder, or psychotic disorde NOS.
Interventions	1. CBT 2. Befriending 3. Routine care
Outcomes	Social Functioning Scale (SFS), Positive and Negative Syndrome Scale (PANSS), Psychotic Symptom Rating Scales (PSYRATS), Calgary Depression Scale for Schizophrenia (CDSS) The Time-Line Follow Back (TLFB), Alcohol and Drug Use Scale (AUS; DUS), Medication Event Monitoring System (MEMS), Rosenberg Self-Esteem Scale, Maastrich Assessment of Coping Skills (MACS).
Starting date	June 2007.
Contact information	Jean Addington, PhD 416-535-8501 Ext. 4360 Jean_Addington@camh.net
Notes	

NCT 2012

Trial name or title	Effectiveness and cost effectiveness of peer mentors in reducing hospital use.
Methods	Randomised controlled trial. Parallel assignment. Open label.
Participants	Diagnosed with serious mental illness.
Interventions	1.Community Connector.



NCT 2012 (Continued)	2. Peer recovery mentor.3. Peer case manager.
Outcomes	Primary Outcome: Service use via self-report from the participant. Two time points: 3 months and 9 months. Secondary Outcome: (measured at 3 months and 9 months) Psychiatric symptoms measured using the Symptom Distress scale, and the Paranoia and Psychoticism subscales from the Symptom Check List-90 (SCL-90),Quality of life measured using Lehman's Brief Quality of Life scale, Community inclusion measured using Mancini's Community Connections Inventory.
Starting date	August 2011.
Contact information	Larry Davidson, PhD 203-764-7583 larry.davidson@yale.edu
Notes	

Robinson 2010

Trial name or title	The development, design, and testing of a peer support intervention in youth being discharged from a first episode psychosis clinic.
Methods	Randomised controlled trial, open-label, parallel group.
Participants	Outpatients diagnosed with psychosis.
Interventions	Peer-support for 6 months: 2 hours of contact per fortnight to engage and support the participant in transition to the new service (accompanying, helping with attendance, understanding health services, and motivating to develop contacts) Treatment as usual
Outcomes	Not reported.
Starting date	Not reported.
Contact information	Jo Robinson, Orygen Youth Health Research Centre, University of Melbourne, Melbourne, Australia
Notes	DATA COLLECTION HAS NOT STARTED

Rohricht 2000

Trial name or title	RCT in effectiveness of body oriented psychotherapy on anergia in patients with chronic schizophrenia.	
Methods		
Participants	People with schizophrenia.	
Interventions	Body oriented psychotherapy. No information available for comparison group.	



Rohricht 2000 (Continued)	
Outcomes	
Starting date	
Contact information	
Notes	
Sensky 2000a	
Trial name or title	A five-year follow-up of patients with schizophrenia with drug-resistant symptoms treated with cognitive behaviour therapy or a befriending intervention.
Methods	
Participants	People with schizophrenia.
Interventions	CBT. Befriending intervention.
Outcomes	Global state: Comprehensive Psychopathological Rating Scale. Mental state: Montgomery-Asberg Depression Rating Score; the Schedule for Negative Symptoms.
Starting date	
Contact information	
Notes	
Tarrier 2000c	
Trial name or title	A psychological intervention programme to reduce positive symptoms and prevent relapse in psychotic patients.
Methods	
Participants	People with schizophrenia.
Interventions	 Psychological intervention. Supportive counselling and routine care.
Outcomes	Relapse and social functioning.
Starting date	
Contact information	
Notes	



ICD - International Classification of Diseases RCT - randomised controlled trial

DATA AND ANALYSES

Comparison 1. SUPPORTIVE THERAPY versus STANDARD CARE

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
1 Global state: Relapse	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only	
1.1 medium term	1	54	Risk Ratio (M-H, Fixed, 95% CI)	0.12 [0.01, 2.11]	
1.2 long term	1	54	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.44, 2.11]	
2 Service outcomes: Hospitalisation	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only	
2.1 long term	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.07, 15.08]	
3 Mental state. 1. No clinically important improvement in general mental state	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only	
3.1 medium term	1	54	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.77, 1.17]	
3.2 long term	2	98	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.82, 1.11]	
4 Mental state: 2. Average endpoint general mental state score (PANSS, high = poor)	2		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	
4.1 short term	1	131	Mean Difference (IV, Fixed, 95% CI)	-4.42 [-10.13, 1.29]	
4.2 long term	1	36	Mean Difference (IV, Fixed, 95% CI)	4.70 [-6.71, 16.11]	
5 Mental state: 3. Average endpoint negative symptoms score (SANS, skewed data, high = poor)			Other data	No numeric data	
5.1 short term			Other data	No numeric data	
5.2 long term	,		Other data	No numeric data	
6 Mental state: 4. Average endpoint positive symptoms score (PANSS positive subscale, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
6.1 short term	1	131	Mean Difference (IV, Fixed, 95% CI)	-1.09 [-2.84, 0.66]	
7 Mental state: 5. Average endpoint positive symptoms score (BPRS, skewed data, high = poor)			Other data	No numeric data	
7.1 short term			Other data	No numeric data	
7.2 long term			Other data	No numeric data	
8 Mental state: 6. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)			Other data	No numeric data	
8.1 short term			Other data	No numeric data	
8.2 long term			Other data	No numeric data	
9 Mental state: 7. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)			Other data	No numeric data	
9.1 long term			Other data	No numeric data	
10 Mental state: 8. Average endpoint depressive symptoms score (CESD, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	
10.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	1.61 [-1.61, 4.83]	
11 Leaving the study early	4	354	Risk Ratio (M-H, Fixed, 95% CI)	0.86 [0.53, 1.40]	
12 General functioning: 2. Average endpoint general functioning score (GAF-M, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	
12.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	-2.66 [-6.20, 0.88]	
13 General functioning: 1. Average endpoint general functioning score (GAS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	
13.1 long term	1	29	Mean Difference (IV, Fixed, 95% CI)	1.40 [-5.09, 7.89]	
14 General functioning: 3. Average endpoint social functioning score (SFS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only	

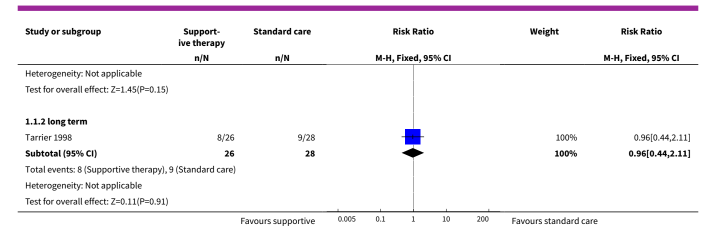


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
14.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	-0.67 [-7.05, 5.71]
15 Satisfaction with treatment: Recipient of care not satisfied with treatment	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
15.1 long term	1	44	Risk Ratio (M-H, Fixed, 95% CI)	1.83 [0.75, 4.47]
16 Quality of life: 1. Average endpoint self-esteem score (RSES, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
16.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	-1.21 [-2.85, 0.43]
17 Quality of life: 2. Average endpoint well-being score (WBS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
17.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	-2.73 [-6.04, 0.58]
18 Quality of life: 3. Average endpoint global health score (GHQ, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
18.1 long term	1	260	Mean Difference (IV, Fixed, 95% CI)	2.45 [-2.41, 7.31]
19 Death	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
19.1 medium term	1	54	Risk Ratio (M-H, Fixed, 95% CI)	3.22 [0.14, 75.75]
19.2 long term	2	92	Risk Ratio (M-H, Fixed, 95% CI)	2.87 [0.31, 26.63]
20 Medication: No reduction in dose of antipsychotic medication	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
20.1 long term	1	44	Risk Ratio (M-H, Fixed, 95% CI)	0.81 [0.59, 1.12]

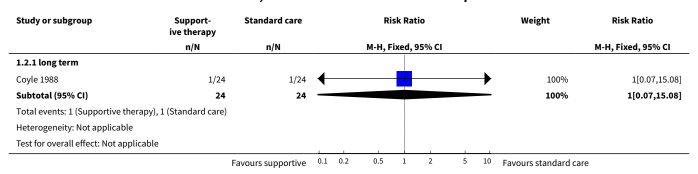
Analysis 1.1. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 1 Global state: Relapse.

Study or subgroup	Support- ive therapy			F	lisk Rati	o		Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 9	5% CI			M-H, Fixed, 95% CI
1.1.1 medium term									
Tarrier 1998	0/26	4/28		-	_			100%	0.12[0.01,2.11]
Subtotal (95% CI)	26	28			-			100%	0.12[0.01,2.11]
Total events: 0 (Supportive th	erapy), 4 (Standard care)								
	ī	Favours supportive	0.005	0.1	1	10	200	Favours standard care	





Analysis 1.2. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 2 Service outcomes: Hospitalisation.

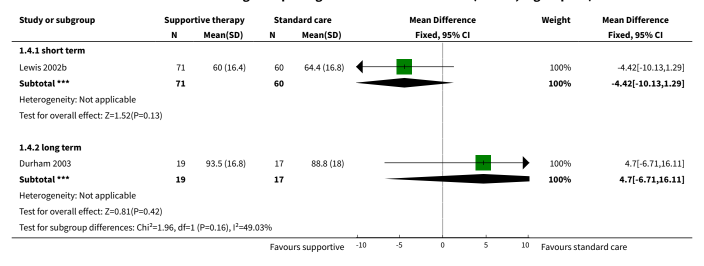


Analysis 1.3. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 3 Mental state. 1. No clinically important improvement in general mental state.

Study or subgroup	Support- ive therapy	Standard care		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		М-Н,	Fixed, 95%	CI			M-H, Fixed, 95% CI
1.3.1 medium term									
Tarrier 1998	22/26	25/28			+			100%	0.95[0.77,1.17]
Subtotal (95% CI)	26	28			•			100%	0.95[0.77,1.17]
Total events: 22 (Supportive therap	y), 25 (Standard care)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.51(P=0.6	1)								
1.3.2 long term									
Durham 2003	20/23	19/21			•			45.21%	0.96[0.78,1.19]
Tarrier 1998	22/26	25/28			•			54.79%	0.95[0.77,1.17]
Subtotal (95% CI)	49	49			♦			100%	0.95[0.82,1.11]
Total events: 42 (Supportive therap	y), 44 (Standard care)								
Heterogeneity: Tau ² =0; Chi ² =0.01, d	f=1(P=0.93); I ² =0%								
Test for overall effect: Z=0.62(P=0.5	3)								
	F	avours supportive	0.005	0.1	1	10	200	Favours standard care	



Analysis 1.4. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 4 Mental state: 2. Average endpoint general mental state score (PANSS, high = poor).



Analysis 1.5. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 5 Mental state: 3. Average endpoint negative symptoms score (SANS, skewed data, high = poor).

Mental state: 3. Average endpoint negative symptoms score (SANS, skewed data, high = poor)

Study	Intervention	N	Mear	n SD	
		short to	erm		
Tarrier 1998	Supportive therapy	21	10.19	5.83	
Tarrier 1998	Standard care	26	10.73	4.19	
		long te	rm		
Tarrier 1998	Supportive therapy	21	9.90	5.44	
Tarrier 1998	Standard care	26	11.46	4.34	

Analysis 1.6. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 6 Mental state: 4. Average endpoint positive symptoms score (PANSS positive subscale, high = poor).

Study or subgroup	Supportive therapy		Stan	Standard care		Mean Difference				Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		ı	Fixed, 95% CI				Fixed, 95% CI
1.6.1 short term											
Lewis 2002b	71	12.6 (4.8)	60	13.7 (5.3)			-			100%	-1.09[-2.84,0.66]
Subtotal ***	71		60							100%	-1.09[-2.84,0.66]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.22(P=0.22)										
			Favou	rs supportive	-10	-5	0	5	10	Favours sta	ndard care



Analysis 1.7. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 7 Mental state: 5. Average endpoint positive symptoms score (BPRS, skewed data, high = poor).

Mental state: 5. Average endpoint positive symptoms score (BPRS, skewed data, high = poor)

Study	Intervention	N	Mean	SD								
	short term											
Tarrier 1998	Supportive therapy	21	15.81	16.10								
Tarrier 1998	Standard care	26	15.65	14.62								
	long term											
Davidson 2004	Supportive therapy	190	10.71	7.92								
Davidson 2004	Standard care	70	9.14	6.92								
Tarrier 1998	Supportive therapy	21	16.30	16.71								
Tarrier 1998	Standard care	26	17.63	13.51								

Analysis 1.8. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 8 Mental state: 6. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor).

Mental state: 6. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)

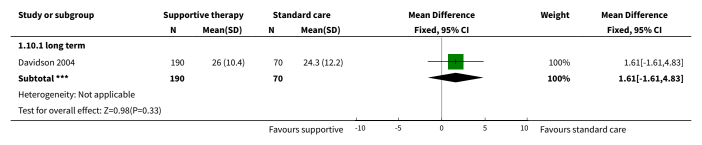
Study	Intervention	Mean	SD	N	
		short ter	m		
Lewis 2002b	Supportive therapy	6.13	6.98	67	
Lewis 2002b	Standard care	7.52	7.15	56	
		long terr	n		
Durham 2003	Supportive therapy	9.7	6.1	19	
Durham 2003	Standard care	11.2	6.5	18	

Analysis 1.9. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 9 Mental state: 7. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor).

Mental state: 7. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)

Study	Intervention	Mean	SD		N
		long term	1		
Durham 2003	Supportive therapy	18.0	12.2	19	
Durham 2003	Standard care	17.2	11.7	17	

Analysis 1.10. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 10 Mental state: 8. Average endpoint depressive symptoms score (CESD, high = poor).





Analysis 1.11. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 11 Leaving the study early.

Study or subgroup	Support- ive therapy	Standard care	ard care Risk Ratio			Weight	Risk Ratio		
	n/N	n/N		М-Н, Е	ixed, 9	5% CI			M-H, Fixed, 95% CI
Coyle 1988	3/24	3/24		_	_	_		10.15%	1[0.22,4.47]
Durham 2003	4/23	4/21		-	+	-		14.15%	0.91[0.26,3.2]
Lewis 2002b	18/106	21/102						72.44%	0.82[0.47,1.46]
Tarrier 1998	1/26	1/28			\rightarrow			3.26%	1.08[0.07,16.35]
Total (95% CI)	179	175			•			100%	0.86[0.53,1.4]
Total events: 26 (Supportive t	herapy), 29 (Standard care)							
Heterogeneity: Tau ² =0; Chi ² =0	0.09, df=3(P=0.99); I ² =0%								
Test for overall effect: Z=0.6(P	=0.55)								
	-	avours supportive	0.001	0.1	1	10	1000	Favours standard care	

Analysis 1.12. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 12 General functioning: 2. Average endpoint general functioning score (GAF-M, high = good).

Study or subgroup	Supportive therapy		Stan	Standard care		Mean Difference				Weight	Mean Difference Fixed, 95% CI	
	N	Mean(SD)	N Mean(SD)		Fixed, 95% CI							
1.12.1 long term										,		
Davidson 2004	190	49.9 (11.6)	70	52.6 (13.4)						100%	-2.66[-6.2,0.88]	
Subtotal ***	190		70							100%	-2.66[-6.2,0.88]	
Heterogeneity: Not applicable												
Test for overall effect: Z=1.47(P=0.14	.)											
			Favours	standard care	-10	-5	0	5	10	Favours supp	ortive	

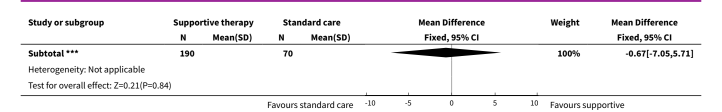
Analysis 1.13. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 13 General functioning: 1. Average endpoint general functioning score (GAS, high = good).

Study or subgroup	Suppor	tive therapy	Stan	dard care		Mea	n Differen	ce		Weight I	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ed, 95% C	ı			Fixed, 95% CI
1.13.1 long term											
Durham 2003	12	36.3 (9.8)	17	34.9 (7.1)			-		_	100%	1.4[-5.09,7.89]
Subtotal ***	12		17						-	100%	1.4[-5.09,7.89]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.42(P=0.67	')										
			Favours	standard care	-10	-5	0	5	10	Favours support	ive

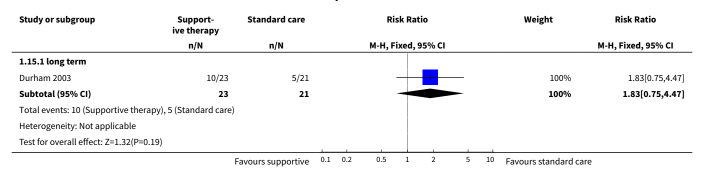
Analysis 1.14. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 14 General functioning: 3. Average endpoint social functioning score (SFS, high = good).

Study or subgroup	Suppor	tive therapy	Star	ndard care		Me	an Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95%	CI			Fixed, 95% CI
1.14.1 long term											
Davidson 2004	190	126.6 (23.4)	70	127.3 (23.3)						100%	-0.67[-7.05,5.71]
			Favours	standard care	-10	-5	0	5	10	Favours supp	portive

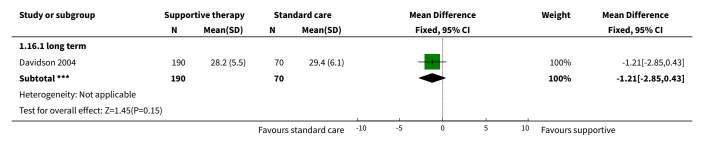




Analysis 1.15. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 15 Satisfaction with treatment: Recipient of care not satisfied with treatment.



Analysis 1.16. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 16 Quality of life: 1. Average endpoint self-esteem score (RSES, high = good).



Analysis 1.17. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 17 Quality of life: 2. Average endpoint well-being score (WBS, high = good).

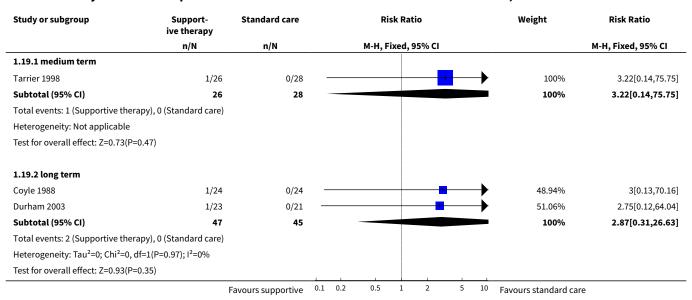
Study or subgroup	Suppor	tive therapy	Stan	dard care		Mea	n Differer	ice		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ed, 95% (CI			Fixed, 95% CI
1.17.1 long term											
Davidson 2004	190	56.5 (11.1)	70	59.2 (12.4)			<u> </u>			100%	-2.73[-6.04,0.58]
Subtotal ***	190		70							100%	-2.73[-6.04,0.58]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.61(P=0.1	1)										
			Favours	standard care	-10	-5	0	5	10	Favours sup	portive



Analysis 1.18. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 18 Quality of life: 3. Average endpoint global health score (GHQ, high = poor).

Study or subgroup	Suppor	tive therapy	Stan	dard care		Me	ean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	Fixed, 95% CI			Fixed, 95% CI
1.18.1 long term										
Davidson 2004	190	63.7 (15.6)	70	61.3 (18.5)				_	100%	2.45[-2.41,7.31]
Subtotal ***	190		70					-	100%	2.45[-2.41,7.31]
Heterogeneity: Not applicable										
Test for overall effect: Z=0.99(P=0.32)									
			Favou	rs supportive	-10	-5	0 5	10	Favours star	ndard care

Analysis 1.19. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 19 Death.



Analysis 1.20. Comparison 1 SUPPORTIVE THERAPY versus STANDARD CARE, Outcome 20 Medication: No reduction in dose of antipsychotic medication.

Study or subgroup	Support- ive therapy	Standard care			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
1.20.1 long term											
Durham 2003	16/23	18/21			-	+				100%	0.81[0.59,1.12]
Subtotal (95% CI)	23	21			•					100%	0.81[0.59,1.12]
Total events: 16 (Supportive therap	y), 18 (Standard care))									
Heterogeneity: Not applicable											
Test for overall effect: Z=1.27(P=0.2)											
	F	avours supportive	0.1	0.2	0.5	1	2	5	10	Favours standard care	



Comparison 2. SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: 1. Relapse	6		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
1.1 medium term	2	100	Risk Ratio (M-H, Random, 95% CI)	2.86 [0.32, 25.24]
1.2 long term	5	270	Risk Ratio (M-H, Random, 95% CI)	1.19 [0.66, 2.16]
2 Global state: 2. No remission	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.87 [1.11, 3.15]
3 Service outcomes: Hospitalisation	6		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 medium term	3	153	Risk Ratio (M-H, Fixed, 95% CI)	1.60 [0.85, 3.00]
3.2 long term	4	306	Risk Ratio (M-H, Fixed, 95% CI)	1.82 [1.11, 2.99]
4 Mental state: 1. No clinically important improvement in general mental state	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.1 medium term	1	59	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.95, 1.70]
4.2 long term	3	194	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [1.04, 1.54]
5 Mental state: 2. Average endpoint general mental state score (PANSS, high = poor)	4		Mean Difference (IV, Random, 95% CI)	Totals not selected
5.1 short term	3		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 long term	2		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 Mental state: 3. Average endpoint general mental state score (BPRS, high = poor)	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
6.1 short term	2	92	Mean Difference (IV, Fixed, 95% CI)	-1.07 [-5.08, 2.94]
6.2 medium term	1	37	Mean Difference (IV, Fixed, 95% CI)	7.60 [0.90, 14.30]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
7 Mental state: 4. Average endpoint general mental state score (BPRS short form, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
7.1 short term	1	74	Mean Difference (IV, Fixed, 95% CI)	-0.90 [-3.02, 1.22]
7.2 medium term	1	67	Mean Difference (IV, Fixed, 95% CI)	2.20 [-1.18, 5.58]
7.3 long term	1	45	Mean Difference (IV, Fixed, 95% CI)	2.30 [-0.54, 5.14]
8 Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor)			Other data	No numeric data
8.1 medium term	,		Other data	No numeric data
8.2 long term			Other data	No numeric data
9 Mental state: 6. Average endpoint general mental state score (PANSS general subscale, high = poor)	3		Mean Difference (IV, Random, 95% CI)	Totals not selected
9.1 short term	2		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 medium term	1		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
9.3 long term	1		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
10 Mental state: 7. No clinically important improvement in negative symptoms	1	90	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.63, 1.46]
11 Mental state: 8. No clinically important improvement in depressive symptoms	1	90	Risk Ratio (M-H, Fixed, 95% CI)	1.53 [0.92, 2.55]
12 Mental state: 9. Episode of affective symptoms	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
12.1 long term	2	151	Risk Ratio (M-H, Fixed, 95% CI)	1.84 [1.15, 2.94]
13 Mental state: 10. Average endpoint negative symptoms score (PANSS negative, high=poor)	3		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
13.1 short term	2		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
13.2 medium term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
13.3 long term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
14 Mental state: 11. Average endpoint negative symptoms score (SANS, high=poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
14.1 medium term	1	37	Mean Difference (IV, Fixed, 95% CI)	6.60 [-5.81, 19.01]
15 Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor)			Other data	No numeric data
15.1 short term			Other data	No numeric data
15.2 medium term			Other data	No numeric data
15.3 long term			Other data	No numeric data
16 Mental state: 13. Average endpoint positive symptoms score (PANSS positive subscale, high = poor)	4		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
16.1 short term	3		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
16.2 medium term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
16.3 long term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
17 Mental state: 14. Average end- point positive symptoms score (SAPS, skewed data, high = poor)			Other data	No numeric data
17.1 medium term			Other data	No numeric data
18 Mental state: 15. Average endpoint positive symptoms score (BPRS, skewed data, high = poor)			Other data	No numeric data
18.1 short term	,		Other data	No numeric data
18.2 long term			Other data	No numeric data
19 Mental state: 16. Average endpoint score (PANSS thought disturbance cluster, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
19.1 short term	1	12	Mean Difference (IV, Fixed, 95% CI)	4.3 [1.17, 7.43]
20 Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)			Other data	No numeric data
20.1 short term			Other data	No numeric data
20.2 long term			Other data	No numeric data
21 Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)			Other data	No numeric data
21.1 long term			Other data	No numeric data
22 Mental state: 19. Average endpoint voices score (PSYRATS voices score, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
22.1 short term	1	65	Mean Difference (IV, Fixed, 95% CI)	0.10 [-3.63, 3.83]
22.2 long term	1	65	Mean Difference (IV, Fixed, 95% CI)	0.0 [-4.76, 4.76]
23 Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data)			Other data	No numeric data
23.1 Malevolence			Other data	No numeric data
23.2 Benevolence			Other data	No numeric data
23.3 Resistance			Other data	No numeric data
23.4 Engagement			Other data	No numeric data
23.5 Omnipotence			Other data	No numeric data
24 Mental state: 21. Average endpoint depression score (MADRS, skewed da- ta, high = poor)			Other data	No numeric data
24.1 medium term			Other data	No numeric data
24.2 long term	,		Other data	No numeric data
25 Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor)			Other data	No numeric data



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
26 Leaving the study early	19	1412	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.87, 1.21]
26.1 general reasons	17	1261	Risk Ratio (M-H, Fixed, 95% CI)	0.97 [0.83, 1.15]
26.2 treatment-related reasons	2	151	Risk Ratio (M-H, Fixed, 95% CI)	2.15 [1.07, 4.31]
27 General functioning: 1. Average endpoint general functioning score (GAF & GAS, high = good)	2		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
27.1 short term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
27.2 medium term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
27.3 long term	2		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
28 General functioning: 2. Average endpoint social functioning score (SFS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
28.1 short term	1	65	Mean Difference (IV, Fixed, 95% CI)	-7.20 [-17.86, 3.46]
28.2 long term	1	65	Mean Difference (IV, Fixed, 95% CI)	-8.80 [-21.67, 4.07]
29 General functioning: 3. No paid work	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
29.1 long term	2	171	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.84, 1.25]
30 General functioning: 4. Admission to residential placement	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
30.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.24, 4.59]
31 General functioning: 5. Admission to jail	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
31.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.24, 4.59]
32 Satisfaction with treatment: Recipient of care not satisfied with treatment	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
32.1 long term	1	45	Risk Ratio (M-H, Fixed, 95% CI)	3.19 [1.01, 10.07]
33 Quality of life: Average endpoint quality of life score (QLS, high = good)	1	19	Mean Difference (IV, Fixed, 95% CI)	-0.07 [-21.11, 20.97]
34 Death	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
34.1 medium term	1	59	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.08, 19.34]
34.2 long term	2	115	Risk Ratio (M-H, Fixed, 95% CI)	3.99 [0.44, 36.08]
35 Behaviour: 1. Social impairment on SBAS	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
35.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.46 [1.04, 2.04]
36 Behaviour: 2. Poor coping style with relatives	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
36.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.37, 2.20]
37 Engagement with services: Poor attendance at appointments	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.93 [0.89, 4.17]
38 Insight: Average endpoint insight score (SAI, high = good, skewed data)			Other data	No numeric data
38.1 short term			Other data	No numeric data
38.2 medium term			Other data	No numeric data
38.3 long term			Other data	No numeric data
39 Compliance: 1. Poor compliance to therapy	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
39.1 medium term	2	58	Risk Ratio (M-H, Fixed, 95% CI)	2.63 [1.30, 5.35]
39.2 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.29 [0.69, 2.39]
40 Compliance: 2. adherence to medication (self-report)	1	19	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.83, 1.21]
41 Compliance: 3. average endpoint adherence score (ROMI, high = good)	1	19	Mean Difference (IV, Fixed, 95% CI)	0.45 [-3.58, 4.48]

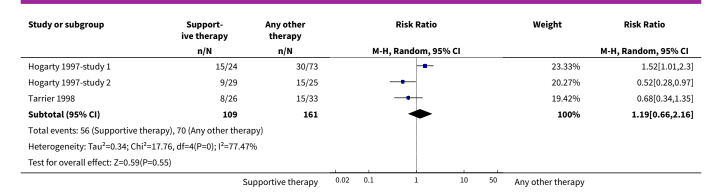


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
42 Compliance: 4. average endpoint non-adherence score (ROMI, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	1.75 [-0.61, 4.11]
43 Medication: Prescribed IM depot medication	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
43.1 medium term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	2.11 [0.99, 4.47]
44 Attitude to medication: 1. Average endpoint attitude to medication score (AMQ, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
44.1 short term	1	74	Mean Difference (IV, Fixed, 95% CI)	-4.50 [-6.83, -2.17]
45 Attitude to medication: 2. Average endpoint attitude to medication score (DAI, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
45.1 short term	1	63	Mean Difference (IV, Fixed, 95% CI)	-5.70 [-9.35, -2.05]
45.2 long term	1	44	Mean Difference (IV, Fixed, 95% CI)	-4.90 [-9.38, -0.42]
46 Economic outcomes: Direct costs (skewed data, not ITT)			Other data	No numeric data
46.1 medium term			Other data	No numeric data
46.2 long term			Other data	No numeric data

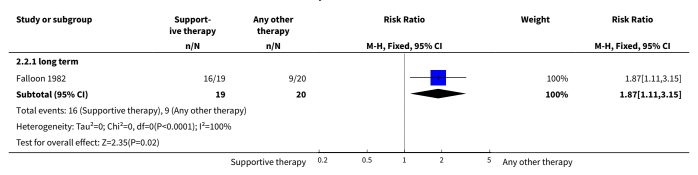
Analysis 2.1. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 1 Global state: 1. Relapse.

Study or subgroup	Support- ive therapy	Any other therapy	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Random, 95% CI		M-H, Random, 95% CI
2.1.1 medium term					
Pinto 1999	3/21	1/20	- 1	100%	2.86[0.32,25.24]
Tarrier 1998	0/26	0/33			Not estimable
Subtotal (95% CI)	47	53		100%	2.86[0.32,25.24]
Total events: 3 (Supportive therapy)	, 1 (Any other therapy)			
Heterogeneity: Not applicable					
Test for overall effect: Z=0.94(P=0.34	.)				
2.1.2 long term					
Falloon 1982	16/19	5/20		18%	3.37[1.54,7.38]
Haddock 1999	8/11	5/10	+	18.97%	1.45[0.71,2.98]
	Sı	pportive therapy	0.02 0.1 1 10 50	Any other therapy	





Analysis 2.2. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 2 Global state: 2. No remission.



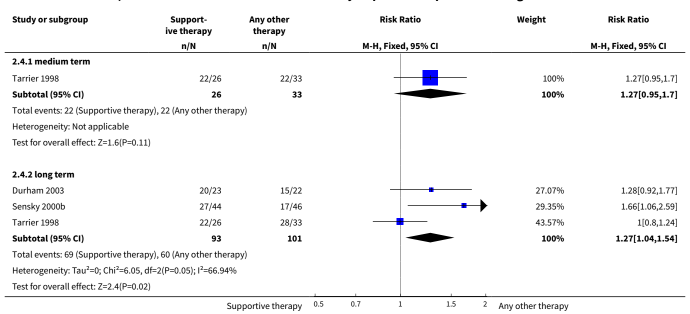
Analysis 2.3. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 3 Service outcomes: Hospitalisation.

Study or subgroup	Support- ive therapy	Any other therapy	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.3.1 medium term					
Kemp 1996	10/22	8/25	- 	64.79%	1.42[0.68,2.95]
Penn 2009	5/33	3/32		26.35%	1.62[0.42,6.21]
Pinto 1999	3/21	1/20	-	8.86%	2.86[0.32,25.24]
Subtotal (95% CI)	76	77		100%	1.6[0.85,3]
Total events: 18 (Supportive t	herapy), 12 (Any other thera	ару)			
Heterogeneity: Tau ² =0; Chi ² =0	0.37, df=2(P=0.83); I ² =0%				
Test for overall effect: Z=1.47(P=0.14)				
2.3.2 long term					
Coyle 1988	1/24	3/46	+	10.85%	0.64[0.07,5.82]
Dincin 1982	19/66	7/66		36.91%	2.71[1.22,6.02]
Falloon 1982	11/19	6/20	-	30.83%	1.93[0.89,4.17]
Penn 2009	3/33	4/32		21.42%	0.73[0.18,3]
Subtotal (95% CI)	142	164	-	100%	1.82[1.11,2.99]
Total events: 34 (Supportive t	herapy), 20 (Any other thera	ару)			
Heterogeneity: Tau ² =0; Chi ² =3	3.46, df=3(P=0.33); I ² =13.41%	6			
	Si	upportive therapy	0.1 0.2 0.5 1 2 5	10 Any other therapy	



Study or subgroup	Support- ive therapy	Any other therapy			Ri	sk Ra	tio			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed,	95% CI				M-H, Fixed, 95% CI
Test for overall effect: Z=2.37(P=0.02)											
		Supportive therapy	0.1	0.2	0.5	1	2	5	10	Any other therapy	

Analysis 2.4. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 4 Mental state: 1. No clinically important improvement in general mental state.

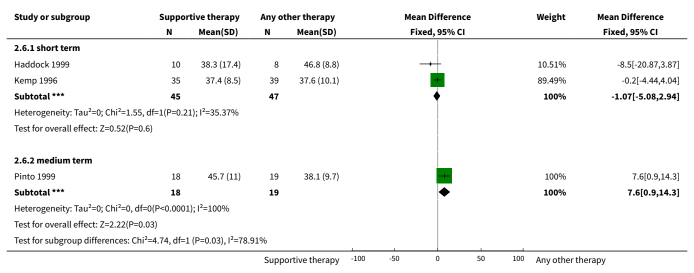


Analysis 2.5. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 5 Mental state: 2. Average endpoint general mental state score (PANSS, high = poor).

Study or subgroup	Suppo	ortive therapy	Any o	other therapy		Меа	an Differe	nce		Mean Difference
	N	Mean(SD)	N	Mean(SD)		Rar	dom, 95%	6 CI		Random, 95% CI
2.5.1 short term										
Levine 1998	6	60.5 (3)	6	29.5 (4.9)				-		31[26.4,35.6]
Lewis 2002b	71	60 (16.4)	78	61.7 (19.7)			+			-1.77[-7.57,4.03]
Penn 2009	33	59.9 (10.5)	32	52.2 (10.7)			-			7.7[2.54,12.86]
2.5.2 long term										
Durham 2003	19	93.5 (16.8)	21	87 (23.1)			++			6.5[-5.94,18.94]
Penn 2009	33	58.4 (11.2)	32	52.7 (10.1)						5.7[0.52,10.88]
			S	upportive therapy	-50	-25	0	25	50	Any other therapy



Analysis 2.6. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 6 Mental state: 3. Average endpoint general mental state score (BPRS, high = poor).



Analysis 2.7. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 7 Mental state: 4. Average endpoint general mental state score (BPRS short form, high = poor).

Study or subgroup	Suppor	tive therapy	Any ot	her therapy	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.7.1 short term						,	
Kemp 1996	35	11.7 (3.3)	39	12.6 (5.8)		100%	-0.9[-3.02,1.22]
Subtotal ***	35		39			100%	-0.9[-3.02,1.22]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.83(P=0.41	1)						
2.7.2 medium term							
Kemp 1996	31	16.7 (6.9)	36	14.5 (7.2)		100%	2.2[-1.18,5.58]
Subtotal ***	31		36			100%	2.2[-1.18,5.58]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.28(P=0.2)							
2.7.3 long term							
Kemp 1996	20	14.8 (4.1)	25	12.5 (5.6)		100%	2.3[-0.54,5.14]
Subtotal ***	20		25			100%	2.3[-0.54,5.14]
Heterogeneity: Tau ² =0; Chi ² =0, df=0	(P<0.0001	L); I ² =100%					
Test for overall effect: Z=1.59(P=0.11	1)						
Test for subgroup differences: Chi ² =	4.15, df=1	(P=0.13), I ² =51.	85%				
			Sunno	ortive therapy -4	-2 0 2	4 Any other tl	nerany



Analysis 2.8. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 8 Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor).

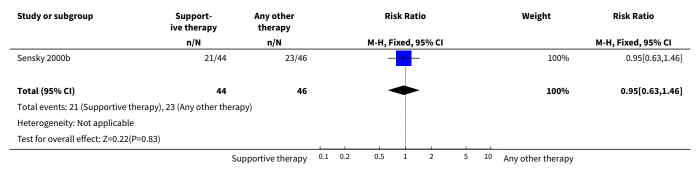
Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor)

Intervention	N	Mean	SD	
	medium ter	m		
Supportive therapy	44	22.9	17.3	
СВТ	46	20.5	13.1	
	long term			
Supportive therapy	44	26.6	25.3	
СВТ	46	15.1	12.0	·
	Supportive therapy CBT Supportive therapy	Supportive therapy 44 CBT 46 Iong term Supportive therapy 44	medium termSupportive therapy4422.9CBT4620.5long termSupportive therapy4426.6	medium term Supportive therapy 44 22.9 17.3 CBT 46 20.5 13.1 long term Supportive therapy 44 26.6 25.3

Analysis 2.9. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 9 Mental state: 6. Average endpoint general mental state score (PANSS general subscale, high = poor).

Study or subgroup	Suppo	ortive therapy	Any o	other therapy	Mean Difference	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI	Random, 95% CI
2.9.1 short term						
Levine 1998	6	31.8 (2.8)	6	14.7 (3.1)		17.1[13.76,20.44]
Penn 2009	33	30 (7.1)	32	25.6 (5.3)		4.4[1.36,7.44]
2.9.2 medium term						
Uzenoff 2007	9	27.6 (8.6)	10	24.7 (3.7)	+-	2.86[-3.21,8.93]
2.9.3 long term						
Penn 2009	33	29.6 (6.2)	32	26 (5.1)		3.6[0.84,6.36]
			S	upportive therapy	-20 -10 0 10 20	Any other therapy

Analysis 2.10. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 10 Mental state: 7. No clinically important improvement in negative symptoms.





Analysis 2.11. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 11 Mental state: 8. No clinically important improvement in depressive symptoms.

Study or subgroup	Support- ive therapy	Any other therapy		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95%	CI			M-H, Fixed, 95% CI
Sensky 2000b	22/44	15/46			•	→	100%	1.53[0.92,2.55]
Total (95% CI)	44	46					100%	1.53[0.92,2.55]
Total events: 22 (Supportive the	erapy), 15 (Any other thera	oy)						
Heterogeneity: Not applicable								
Test for overall effect: Z=1.64(P=	=0.1)							
	Su	pportive therapy	0.5	0.7 1	1.5	2	Any other therapy	

Analysis 2.12. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 12 Mental state: 9. Episode of affective symptoms.

Study or subgroup	Support- ive therapy	Any other therapy			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			М-Н, F	ixed,	95% CI				M-H, Fixed, 95% CI
2.12.1 long term											
Hogarty 1997-study 1	12/24	19/73				-	-			59.33%	1.92[1.1,3.35]
Hogarty 1997-study 2	12/29	6/25				+	-	_		40.67%	1.72[0.76,3.92]
Subtotal (95% CI)	53	98				-	•			100%	1.84[1.15,2.94]
Total events: 24 (Supportive the	erapy), 25 (Any other thera	py)									
Heterogeneity: Tau ² =0; Chi ² =0.0	95, df=1(P=0.83); I ² =0%										
Test for overall effect: Z=2.56(P=	=0.01)										
	Su	pportive therapy	0.1	0.2	0.5	1	2	5	10	Any other therapy	

Analysis 2.13. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 13 Mental state: 10. Average endpoint negative symptoms score (PANSS negative, high=poor).

Study or subgroup	Suppo	ortive therapy	Any o	other therapy		Mean Difference	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed, 95% CI	Fixed, 95% CI
2.13.1 short term							
Levine 1998	6	15 (1.5)	6	7.8 (4)			7.2[3.78,10.62]
Penn 2009	33	12.7 (3.7)	32	12.4 (3.9)		+	0.3[-1.55,2.15]
2.13.2 medium term							
Uzenoff 2007	9	16 (7.4)	10	14.3 (4.9)			1.7[-4,7.4]
2.13.3 long term							
Penn 2009	33	13.2 (4.9)	32	12.9 (4.4)			0.3[-1.96,2.56]
			S	upportive therapy	-10	-5 0 5 10	Any other therapy



Analysis 2.14. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 14 Mental state: 11. Average endpoint negative symptoms score (SANS, high=poor).

Study or subgroup	Suppor	tive therapy	Any ot	her therapy		Mea	n Difference	•		Weight N	lean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ed, 95% CI				Fixed, 95% CI
2.14.1 medium term											
Pinto 1999	18	53.5 (19.1)	19	46.9 (19.4)				-	\rightarrow	100%	6.6[-5.81,19.01]
Subtotal ***	18		19							100%	6.6[-5.81,19.01]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.04(P=0.3))										
			Suppo	rtive therapy	-10	-5	0	5	10	Any other therap	у

Analysis 2.15. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 15 Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor).

Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor)

				,	
Study	Intervention	N	Mean	SD	
		short term			
Tarrier 1998	Supportive therapy	21	10.19	5.48	
Tarrier 1998	СВТ	24	9.83	4.43	
		medium terr	n		
Sensky 2000b	Supportive therapy	44	20.7	20.9	
Sensky 2000b	CBT	46	22.0	17.0	
		long term			
Sensky 2000b	Supportive therapy	44	25.1	27.5	
Sensky 2000b	СВТ	46	18.2	17.7	
Tarrier 1998	Supportive therapy	21	9.90	5.11	
Tarrier 1998	СВТ	23	10.39	3.79	

Analysis 2.16. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 16 Mental state: 13. Average endpoint positive symptoms score (PANSS positive subscale, high = poor).

Study or subgroup	Suppo	ortive therapy	Any o	other therapy	Mean Difference	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	Fixed, 95% CI
2.16.1 short term			,			
Levine 1998	6	13.7 (2.5)	6	7 (4.6)		6.7[2.51,10.89]
Lewis 2002b	71	12.6 (4.8)	78	13 (5.1)	-+	-0.45[-2.03,1.13]
Penn 2009	33	16.5 (4)	32	14.2 (4)		2.3[0.35,4.25]
2.16.2 medium term						
Uzenoff 2007	9	12.1 (4.6)	10	9.4 (2.7)	+-	2.71[-0.71,6.13]
2.16.3 long term						
Penn 2009	33	15.9 (3.6)	32	13.6 (3.4)		2.3[0.6,4]
			S	upportive therapy	-10 -5 0 5 10	Any other therapy



Analysis 2.17. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 17 Mental state: 14. Average endpoint positive symptoms score (SAPS, skewed data, high = poor).

Mental state: 14. Average endpoint positive symptoms score (SAPS, skewed data, high = poor)

Study	Intervention	Mea	n SD	1	N	
		medium	term			
Pinto 1999	Supportive therapy	29.9	12.1	18		
Pinto 1999	CBT	17.9	17.0	19		

Analysis 2.18. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 18 Mental state: 15. Average endpoint positive symptoms score (BPRS, skewed data, high = poor).

Mental state: 15. Average endpoint positive symptoms score (BPRS, skewed data, high = poor)

Study	Intervention	N	Mean	SD				
		short terr	n					
Tarrier 1998	Supportive therapy	21	15.81	16.10				
Tarrier 1998	CBT	23	10.67	9.42				
		long tern	1					
Tarrier 1998	Supportive therapy	21	16.30	16.71				
Tarrier 1998	CBT	23	10.59	11.10				

Analysis 2.19. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 19 Mental state: 16. Average endpoint score (PANSS thought disturbance cluster, high = poor).

Study or subgroup	Suppor	Supportive therapy Any other therapy		Any other therapy Mean Diffe		Mean Difference			Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	xed, 95% CI			Fixed, 95% CI
2.19.1 short term										
Levine 1998	6	7.8 (2.1)	6	3.5 (3.3)				-	100%	4.3[1.17,7.43]
Subtotal ***	6		6					-	100%	4.3[1.17,7.43]
Heterogeneity: Not applicable										
Test for overall effect: Z=2.69(P=0.01)									
			Suppo	rtive therapy	-10	-5	0 5	10	Any other thera	DV

Analysis 2.20. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 20 Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor).

Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)

Study	Intervention	Mean	SD	N	
		short term			
Lewis 2002b	Supportive therapy	6.13	6.98	67	
Lewis 2002b	CBT	6.95	7.66	74	
		long term			
Durham 2003	Supportive therapy	9.7	6.1	19	
Durham 2003	CBT	11.1	5.8	21	

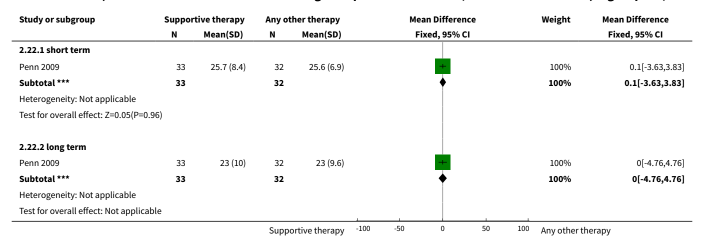


Analysis 2.21. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 21 Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor).

Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)

Study	Intervention	Mean	SD		N	
		long term				
Durham 2003	Supportive therapy	18.0	12.2	19		
Durham 2003	СВТ	18.5	12.8	20		

Analysis 2.22. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 22 Mental state: 19. Average endpoint voices score (PSYRATS voices score, high = poor).



Analysis 2.23. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 23 Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data).

Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data)

Study	Follow-up length	Intervention	Mean	SD	N
		Mal	evolence		
Penn 2009	Short term	Supportive therapy	8.1	5.4	33
Penn 2009	Short term	CBT	6.3	5.4	32
Penn 2009	Long term	Supportive therapy	6.7	4.9	33
Penn 2009	Long term	CBT	6.3	5.4	32
		Ben	evolence		
Penn 2009	Short term	Supportive therapy	5.7	5.5	33
Penn 2009	Short term	CBT	4.4	5.3	32
Penn 2009	Long term	Supportive therapy	6.2	5.4	33
Penn 2009	Long term	CBT	5.3	5.2	32
		Re	sistance		
Penn 2009	Short term	Supportive therapy	16.3	6.6	33
Penn 2009	Short term	CBT	15	7.3	32
Penn 2009	Long term	Supportive therapy	15.3	7.3	33
Penn 2009	Long term	CBT	13.9	7.8	32
		Eng	agement		
Penn 2009	Short term	Supportive therapy	7.8	7.4	33
Penn 2009	Short term	CBT	5.4	5.7	32
Penn 2009	Long term	Supportive therapy	7.3	6.8	33



		Average endpoint beliefs al		•	
Study	Follow-up length	Intervention	Mean	SD	N
Penn 2009	Long term	CBT	6.4	7.2	32
		Omi	nipotence		
Penn 2009	Short term	Supportive therapy	8.6	5	33
Penn 2009	Short term	CBT	8	4.2	32
Penn 2009	Long term	Supportive therapy	7.7	4.3	33
Penn 2009	Long term	CBT	7 3	3 3	32

Analysis 2.24. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 24 Mental state: 21. Average endpoint depression score (MADRS, skewed data, high = poor).

		•		• • •	
Study	Intervention	N	1	Mean	SD
		medium te	rm		
Sensky 2000b	Supportive therapy	44	6.0	5.4	
Sensky 2000b	CBT	46	4.8	3.5	
		long tern	n		
Sensky 2000b	Supportive therapy	44	6.7	7.1	
Sensky 2000b	СВТ	46	3.7	3.2	

Analysis 2.25. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 25 Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor).

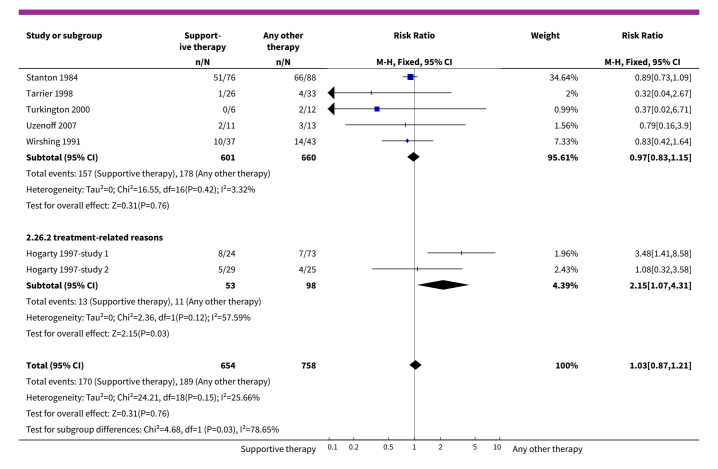
Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor)

Study	Follow-up length	Intervention	Mean	SD	N
Penn 2009	Short term	Supportive therapy	13.9	10.7	33
Penn 2009	Short term	CBT	10.5	8.5	32
Penn 2009	Long term	Supportive therapy	17.9	13.6	33
Penn 2009	Long term	CBT	11.5	9.4	32

Analysis 2.26. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 26 Leaving the study early.

Study or subgroup	Support- ive therapy	Any other therapy	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
2.26.1 general reasons					
Coyle 1988	3/24	10/46		3.88%	0.57[0.17,1.89]
Dincin 1982	29/66	20/66	 	11.33%	1.45[0.92,2.29]
Durham 2003	4/23	1/22	- • -	0.58%	3.83[0.46,31.62]
Eckman 1992	9/21	5/20		2.9%	1.71[0.69,4.24]
Falloon 1982	2/19	3/20		1.66%	0.7[0.13,3.75]
Haddock 1999	0/11	1/10	•	0.89%	0.31[0.01,6.74]
Kemp 1996	15/35	11/39	+	5.89%	1.52[0.81,2.85]
Lewis 2002b	18/106	17/101		9.86%	1.01[0.55,1.85]
Penn 2009	4/33	10/32		5.75%	0.39[0.14,1.11]
Pinto 1999	3/21	1/20	+	0.58%	2.86[0.32,25.24]
Sensky 2000b	6/44	9/46		4.98%	0.7[0.27,1.8]
Spaulding 1999	0/42	1/49	•	0.79%	0.39[0.02,9.27]
	Sı	upportive therapy	0.1 0.2 0.5 1 2 5	¹⁰ Any other therapy	



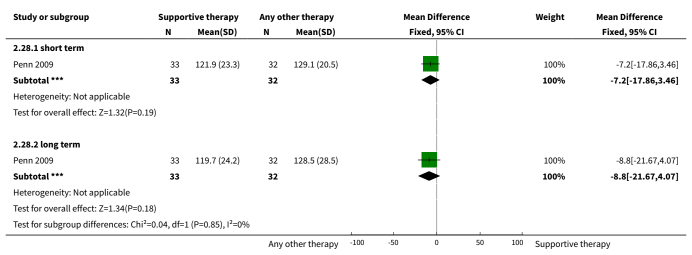


Analysis 2.27. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 27 General functioning: 1. Average endpoint general functioning score (GAF & GAS, high = good).

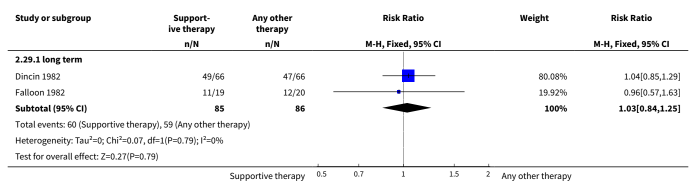
Study or subgroup	Supp	ortive therapy	Any o	other therapy	Mean Difference	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	Fixed, 95% CI
2.27.1 short term						
Kemp 1996	33	44.5 (10.4)	37	54 (17.3)		-9.5[-16.11,-2.89]
2.27.2 medium term						
Kemp 1996	31	43.3 (10.6)	36	55.9 (17.5)		-12.6[-19.43,-5.77]
2.27.3 long term						
Durham 2003	12	36.3 (9.8)	18	35.8 (9.7)		0.5[-6.63,7.63]
Kemp 1996	23	48.3 (14.5)	25	62.8 (18.4)		-14.5[-23.83,-5.17]
				Any other therapy	-20 -10 0 10 20	Supportive therapy



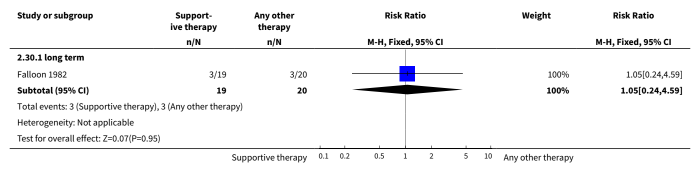
Analysis 2.28. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 28 General functioning: 2. Average endpoint social functioning score (SFS, high = good).



Analysis 2.29. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 29 General functioning: 3. No paid work.

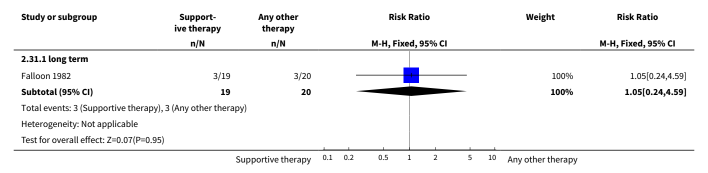


Analysis 2.30. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 30 General functioning: 4. Admission to residential placement.





Analysis 2.31. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 31 General functioning: 5. Admission to jail.



Analysis 2.32. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 32 Satisfaction with treatment: Recipient of care not satisfied with treatment.

Study or subgroup	Support- Any other ive therapy				Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		М-Н	, Fixed, 95%	CI			M-H, Fixed, 95% CI	
2.32.1 long term										
Durham 2003	10/23	3/22					\rightarrow	100%	3.19[1.01,10.07]	
Subtotal (95% CI)	23	22						100%	3.19[1.01,10.07]	
Total events: 10 (Supportive therap	y), 3 (Any other therap	y)								
Heterogeneity: Not applicable										
Test for overall effect: Z=1.98(P=0.0	5)									
	Su	pportive therapy	0.5	0.7	1	1.5	2	Any other therapy		

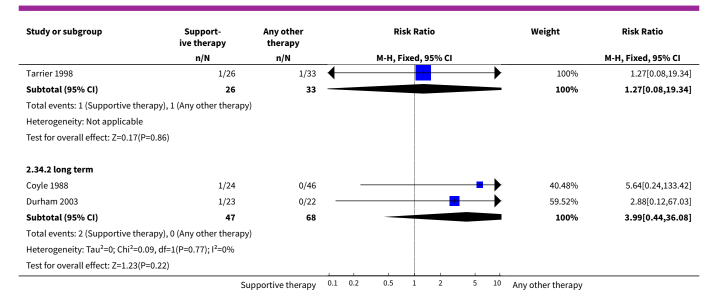
Analysis 2.33. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 33 Quality of life: Average endpoint quality of life score (QLS, high = good).

Study or subgroup	Suppor	tive therapy	Any other therapy			Ме	ean Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	I			Fixed, 95% CI
Uzenoff 2007	9	75.6 (26.6)	10	75.7 (19.1)						100%	-0.07[-21.11,20.97]
Total ***	9		10				•			100%	-0.07[-21.11,20.97]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.01(P=0.9	9)										
			Any	other therapy	-100	-50	0	50	100	Supportive	therapy

Analysis 2.34. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 34 Death.

Study or subgroup	Support- ive therapy	Any other therapy	Risk Ratio					Weight	Risk Ratio		
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
2.34.1 medium term											
		Supportive therapy	0.1	0.2	0.5	1	2	5	10	Any other therapy	





Analysis 2.35. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 35 Behaviour: 1. Social impairment on SBAS.

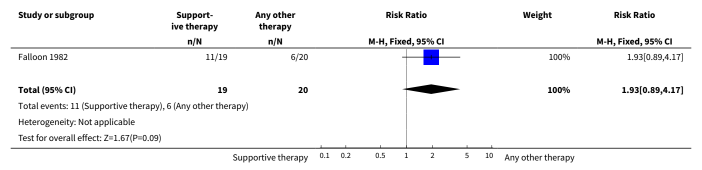
Study or subgroup	Support- ive therapy				Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		М-Н	I, Fixed, 95%	6 CI			M-H, Fixed, 95% CI	
2.35.1 long term										
Falloon 1982	18/19	13/20				-	—	100%	1.46[1.04,2.04]	
Subtotal (95% CI)	19	20			-			100%	1.46[1.04,2.04]	
Total events: 18 (Supportive the	erapy), 13 (Any other thera	oy)								
Heterogeneity: Not applicable										
Test for overall effect: Z=2.18(P=	=0.03)									
	Su	pportive therapy	0.5	0.7	1	1.5	2	Any other therapy		

Analysis 2.36. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 36 Behaviour: 2. Poor coping style with relatives.

Study or subgroup	Support- ive therapy	Any other therapy		Risk Ratio				Weight	Risk Ratio		
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
2.36.1 long term											
Falloon 1982	6/19	7/20				-				100%	0.9[0.37,2.2]
Subtotal (95% CI)	19	20					-			100%	0.9[0.37,2.2]
Total events: 6 (Supportive therapy)	, 7 (Any other therapy))									
Heterogeneity: Not applicable											
Test for overall effect: Z=0.23(P=0.82	2)										
	Su	pportive therapy	0.1	0.2	0.5	1	2	5	10	Any other therapy	



Analysis 2.37. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 37 Engagement with services: Poor attendance at appointments.



Analysis 2.38. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 38 Insight: Average endpoint insight score (SAI, high = good, skewed data).

Insight: Average endpoint insight score (SAI, high = good, skewed data)

		5 ()			
Study	Intervention	Mean	SD		N
		short term			
Kemp 1996	Supportive therapy	40.60	31.20	35	
Kemp 1996	CBT	63.00	23.60	39	
		medium term			
Kemp 1996	Supportive therapy	41.90	30.80	29	
Kemp 1996	СВТ	62.60	23.50	34	
		long term			
Kemp 1996	Supportive therapy	42.60	36.50	20	
Kemp 1996	CBT	63.40	25.50	30	

Analysis 2.39. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 39 Compliance: 1. Poor compliance to therapy.

Study or subgroup	Support- ive therapy	Any other therapy			R	isk Rat	io			Weight	Risk Ratio
	n/N	n/N			М-Н, Г	ixed, 9	5% CI				M-H, Fixed, 95% CI
2.39.1 medium term											
Falloon 1982	15/19	6/20				-	1			100%	2.63[1.3,5.35]
Uzenoff 2007	0/9	0/10									Not estimable
Subtotal (95% CI)	28	30				-		-		100%	2.63[1.3,5.35]
Total events: 15 (Supportive therapy)), 6 (Any other therap	y)									
Heterogeneity: Not applicable											
Test for overall effect: Z=2.68(P=0.01)											
2.39.2 long term											
Falloon 1982	11/19	9/20				-	_			100%	1.29[0.69,2.39]
Subtotal (95% CI)	19	20				\blacktriangleleft	-			100%	1.29[0.69,2.39]
Total events: 11 (Supportive therapy)), 9 (Any other therap	y)									
Heterogeneity: Not applicable											
Test for overall effect: Z=0.8(P=0.42)											
	Sı	pportive therapy	0.1	0.2	0.5	1	2	5	10	Any other therapy	



Analysis 2.40. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 40 Compliance: 2. adherence to medication (self-report).

Study or subgroup	Support- ive therapy	Any other therapy			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-I	H, Fixed, 95% C	:1			M-H, Fixed, 95% CI	
Uzenoff 2007	9/9	10/10			+			100%	1[0.83,1.21]	
Total (95% CI)	9	10			•			100%	1[0.83,1.21]	
Total events: 9 (Supportive therapy),	10 (Any other therap	y)								
Heterogeneity: Not applicable										
Test for overall effect: Not applicable						1				
	Su	pportive therapy	0.01	0.1	1	10	100	Any other therapy		

Analysis 2.41. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 41 Compliance: 3. average endpoint adherence score (ROMI, high = good).

Study or subgroup	Suppor	tive therapy	Any ot	her therapy		Ме	an Differen	ice		Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	:1			Fixed, 95% CI	
Uzenoff 2007	9	16.8 (3.7)	10	16.3 (5.2)			+			100%	0.45[-3.58,4.48]	
Total ***	9		10				•			100%	0.45[-3.58,4.48]	
Heterogeneity: Not applicable												
Test for overall effect: Z=0.22(P=0.83	3)											
			Any	other therapy	-100	-50	0	50	100	Supportive th	nerapy	

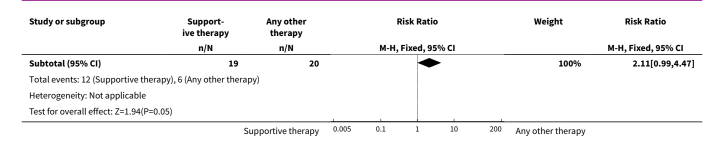
Analysis 2.42. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 42 Compliance: 4. average endpoint non-adherence score (ROMI, high = poor).

Study or subgroup	Suppor	tive therapy	Any other therapy		Mean Difference				Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	I			Fixed, 95% CI
Uzenoff 2007	9	12.1 (2.5)	10	10.4 (2.8)			+			100%	1.75[-0.61,4.11]
Total ***	9		10				•			100%	1.75[-0.61,4.11]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.45(P=0.15	5)										
			Any o	other therapy	-100	-50	0	50	100	Supportive the	ару

Analysis 2.43. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 43 Medication: Prescribed IM depot medication.

Study or subgroup	Support- ive therapy	Any other therapy		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		М-Н,	Fixed, 95	5% CI			M-H, Fixed, 95% CI
2.43.1 medium term									
Falloon 1982	12/19	6/20			-	- ,		100%	2.11[0.99,4.47]
	Sı	pportive therapy	0.005	0.1	1	10	200	Any other therapy	





Analysis 2.44. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 44 Attitude to medication: 1. Average endpoint attitude to medication score (AMQ, high = good).

Study or subgroup	Suppor	tive therapy	Any ot	her therapy		Mea	n Differen	ice		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fix	ed, 95% C	:1			Fixed, 95% CI
2.44.1 short term											
Kemp 1996	35	14.9 (6.1)	39	19.4 (3.7)						100%	-4.5[-6.83,-2.17]
Subtotal ***	35		39							100%	-4.5[-6.83,-2.17]
Heterogeneity: Not applicable											
Test for overall effect: Z=3.78(P=0)											
			Any	other therapy	-10	-5	0	5	10	Supportive t	:herapy

Analysis 2.45. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 45 Attitude to medication: 2. Average endpoint attitude to medication score (DAI, high = good).

Study or subgroup	Suppor	tive therapy	Any ot	her therapy	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
2.45.1 short term							
Kemp 1996	28	44.4 (8.1)	35	50.1 (6.3)		100%	-5.7[-9.35,-2.05]
Subtotal ***	28		35			100%	-5.7[-9.35,-2.05]
Heterogeneity: Not applicable							
Test for overall effect: Z=3.06(P=0)							
2.45.2 long term							
Kemp 1996	16	44.6 (7.5)	28	49.5 (6.9)		100%	-4.9[-9.38,-0.42]
Subtotal ***	16		28			100%	-4.9[-9.38,-0.42]
Heterogeneity: Not applicable							
Test for overall effect: Z=2.15(P=0.0	3)						
Test for subgroup differences: Chi ² =	=0.07, df=1	. (P=0.79), I ² =0%)				
			Any	other therapy	-10 -5 0 5	10 Supportive	therapy

Analysis 2.46. Comparison 2 SUPPORTIVE THERAPY versus ANY OTHER PSYCHOLOGICAL OR PSYCHOSOCIAL TREATMENT, Outcome 46 Economic outcomes: Direct costs (skewed data, not ITT).

Economic outcomes: Direct costs (skewed data, not ITT)

Study	Intervention	Me	an	SD	N
		mediur	n term		
Kemp 1996	Supportive therapy	252	234	34	

CBT

24



Kemp 1996

	Economic outcomes: Direct costs (skewed data, not ITT)									
Study	Intervention	Mean	SD	N						
Kemp 1996	СВТ	187	292	36						
long term										
Kamp 1006	Cupportive thereny	226	404	21						

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Comparison 3. SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: Relapse	5		Risk Ratio (M-H, Fixed, 95% CI)	Totals not selected
1.1 medium term	2		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
1.2 long term	4		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
2 Service outcomes: Hospitalisation	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 medium term	3	153	Risk Ratio (M-H, Fixed, 95% CI)	1.60 [0.85, 3.00]
2.2 long term	1	65	Risk Ratio (M-H, Fixed, 95% CI)	0.73 [0.18, 3.00]
3 Mental state: 1. No clinically important improvement in general mental state	3		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
3.1 medium term	1	59	Risk Ratio (M-H, Random, 95% CI)	1.27 [0.95, 1.70]
3.2 long term	3	194	Risk Ratio (M-H, Random, 95% CI)	1.23 [0.89, 1.70]
4 Mental state: 2. Average endpoint general mental state score (PANSS total, high = poor)	4		Mean Difference (IV, Random, 95% CI)	Totals not selected
4.1 short term	3		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 long term	2		Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Mental state: 3. Average endpoint general mental state score (BPRS, high = poor)	3		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
5.1 short term	2	92	Mean Difference (IV, Fixed, 95% CI)	-1.07 [-5.08, 2.94]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.2 medium term	1	37	Mean Difference (IV, Fixed, 95% CI)	7.60 [0.90, 14.30]
6 Mental state: 4. Average endpoint general mental state score (BPRS short form, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
6.1 short term	1	74	Mean Difference (IV, Fixed, 95% CI)	-0.90 [-3.02, 1.22]
6.2 medium term	1	67	Mean Difference (IV, Fixed, 95% CI)	2.20 [-1.18, 5.58]
6.3 long term	1	45	Mean Difference (IV, Fixed, 95% CI)	2.30 [-0.54, 5.14]
7 Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor)			Other data	No numeric data
7.1 medium term			Other data	No numeric data
7.2 long term			Other data	No numeric data
8 Mental state: 6. Average endpoint general mental state score (PANSS general subscale, high = poor)	2		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
8.1 short term	2		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
8.2 long term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
9 Mental state: 7. No clinically important improvement in negative symptoms	1	90	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.63, 1.46]
10 Mental state: 8. No clinically important improvement in depressive symptoms	1	90	Risk Ratio (M-H, Fixed, 95% CI)	1.53 [0.92, 2.55]
11 Mental state: 9. Episode of affective symptoms	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
11.1 long term	2	101	Risk Ratio (M-H, Fixed, 95% CI)	2.17 [1.16, 4.06]
12 Mental state: 10. Average endpoint negative symptoms score (PANSS negative subscale, high = poor)	2		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
12.1 short term	2		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.2 long term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
13 Mental state: 11. Average endpoint negative symptoms score (SANS, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
13.1 medium term	1	37	Mean Difference (IV, Fixed, 95% CI)	6.60 [-5.81, 19.01]
14 Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor)			Other data	No numeric data
14.1 short term		•	Other data	No numeric data
14.2 medium term			Other data	No numeric data
14.3 long term			Other data	No numeric data
15 Mental state: 13. Average endpoint positive symptoms score (PANSS positive subscale, high = poor)	3		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
15.1 short term	3		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
15.2 long term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
16 Mental state: 14. Average end- point positive symptoms score (SAPS, skewed data, high = poor)			Other data	No numeric data
16.1 medium term			Other data	No numeric data
17 Mental state: 15. Average end- point positive symptoms score (BPRS, skewed data, high = poor)			Other data	No numeric data
17.1 short term			Other data	No numeric data
17.2 long term			Other data	No numeric data
18 Mental state: 16. Average endpoint score (PANSS thought disturbance cluster, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
18.1 short term	1	12	Mean Difference (IV, Fixed, 95% CI)	4.3 [1.17, 7.43]
19 Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)			Other data	No numeric data



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
19.1 short term			Other data	No numeric data
19.2 long term			Other data	No numeric data
20 Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)			Other data	No numeric data
20.1 long term			Other data	No numeric data
21 Mental state: 19. Average endpoint voices score (PSYRATS voices score, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
21.1 short term	1	65	Mean Difference (IV, Fixed, 95% CI)	0.10 [-3.63, 3.83]
21.2 long term	1	65	Mean Difference (IV, Fixed, 95% CI)	0.0 [-4.76, 4.76]
22 Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data)			Other data	No numeric data
22.1 Malevolence			Other data	No numeric data
22.2 Benevolence			Other data	No numeric data
22.3 Resistance			Other data	No numeric data
22.4 Engagement			Other data	No numeric data
22.5 Omnipotence			Other data	No numeric data
23 Mental state: 21. Average endpoint depression score (MADRS, skewed data, high = poor)			Other data	No numeric data
23.1 medium term			Other data	No numeric data
23.2 long term			Other data	No numeric data
24 Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor)			Other data	No numeric data
25 Leaving the study early	12		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
25.1 general reasons	10	711	Risk Ratio (M-H, Fixed, 95% CI)	0.93 [0.66, 1.30]
25.2 treatment-related reasons	2	101	Risk Ratio (M-H, Fixed, 95% CI)	2.34 [0.90, 6.10]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
26 General functioning: 1. Average endpoint general functioning score (GAF & GAS, high = good)	2		Mean Difference (IV, Fixed, 95% CI)	Totals not selected
26.1 short term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
26.2 medium term	1		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
26.3 long term	2		Mean Difference (IV, Fixed, 95% CI)	0.0 [0.0, 0.0]
27 General functioning: 2. Average endpoint social functioning score (SFS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
27.1 short term	1	65	Mean Difference (IV, Fixed, 95% CI)	-7.20 [-17.86, 3.46]
27.2 long term	1	65	Mean Difference (IV, Fixed, 95% CI)	-8.80 [-21.67, 4.07]
28 Satisfaction with treatment: Recipient of care not satisfied with treatment	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
28.1 long term	1	45	Risk Ratio (M-H, Fixed, 95% CI)	3.19 [1.01, 10.07]
29 Quality of life: Average endpoint score (RSES, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
29.1 short term	1	65	Mean Difference (IV, Fixed, 95% CI)	-0.80 [-3.77, 2.17]
29.2 long term	1	65	Mean Difference (IV, Fixed, 95% CI)	-1.70 [-5.19, 1.79]
30 Death	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
30.1 medium term	1	59	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [0.08, 19.34]
30.2 long term	1	45	Risk Ratio (M-H, Fixed, 95% CI)	2.88 [0.12, 67.03]
31 Insight:1. Average endpoint insight score (SAI, skewed data, high = good)			Other data	No numeric data
31.1 short term			Other data	No numeric data
31.2 medium term			Other data	No numeric data
31.3 long term			Other data	No numeric data



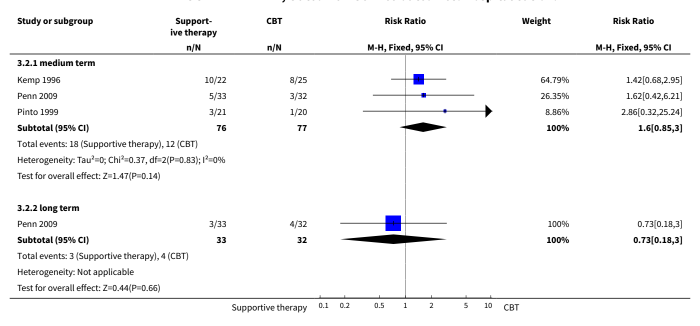
				=
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
32 Insight: 2. Average endpoint insight score (BCIS composite, skewed data, high = good)			Other data	No numeric data
33 Medication: no reduction in dose of antipsychotic medication	1	90	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.68, 1.17]
34 Attitude to medication: 1. Average endpoint attitude to medication score (AMQ, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
34.1 short term	1	74	Mean Difference (IV, Fixed, 95% CI)	-4.50 [-6.83, -2.17]
35 Attitude to medication: 2. Average endpoint attitude to medication score (DAI, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
35.1 short term	1	63	Mean Difference (IV, Fixed, 95% CI)	-5.70 [-9.35, -2.05]
35.2 long term	1	44	Mean Difference (IV, Fixed, 95% CI)	-4.90 [-9.38, -0.42]
36 Economic outcomes: Direct costs (skewed data)			Other data	No numeric data
36.1 medium term			Other data	No numeric data
36.2 long term			Other data	No numeric data

Analysis 3.1. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 1 Global state: Relapse.

Study or subgroup	Supportive therapy	CBT	Risk Ratio	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
3.1.1 medium term				
Pinto 1999	3/21	1/20	- 	2.86[0.32,25.24]
Tarrier 1998	0/26	0/33		Not estimable
3.1.2 long term				
Haddock 1999	8/11	5/10	+-	1.45[0.71,2.98]
Hogarty 1997-study 1	15/24	4/23		3.59[1.4,9.23]
Hogarty 1997-study 2	9/29	15/25		0.52[0.28,0.97]
Tarrier 1998	8/26	15/33	-+	0.68[0.34,1.35]
rarrier 1998	8/26		0.02 0.1 1 10 50	



Analysis 3.2. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 2 Service outcomes: Hospitalisation.



Analysis 3.3. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 3 Mental state: 1. No clinically important improvement in general mental state.

Study or subgroup	Support- ive therapy	СВТ	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Random, 95% CI		M-H, Random, 95% CI	
3.3.1 medium term						
Tarrier 1998	22/26	22/33	 	100%	1.27[0.95,1.7]	
Subtotal (95% CI)	26	33		100%	1.27[0.95,1.7]	
Total events: 22 (Supportive therap	oy), 22 (CBT)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.6(P=0.11	.)					
3.3.2 long term						
Durham 2003	20/23	15/22		33.25%	1.28[0.92,1.77]	
Sensky 2000b	27/44	17/46		25.73%	1.66[1.06,2.59]	
Tarrier 1998	22/26	28/33		41.02%	1[0.8,1.24]	
Subtotal (95% CI)	93	101		100%	1.23[0.89,1.7]	
Total events: 69 (Supportive therap	oy), 60 (CBT)					
Heterogeneity: Tau ² =0.05; Chi ² =6.0	5, df=2(P=0.05); I ² =66.94	%				
Test for overall effect: Z=1.28(P=0.2	2)					
	Sup	portive therapy 0.5	0.7 1 1.5 2	СВТ		



Analysis 3.4. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 4 Mental state: 2. Average endpoint general mental state score (PANSS total, high = poor).

Study or subgroup	subgroup Supportive therapy CBT Mean Difference		Mean Difference	nce Mean Difference		
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI	Random, 95% CI
3.4.1 short term						
Levine 1998	6	60.5 (3)	6	29.5 (4.9)	+	31[26.4,35.6]
Lewis 2002b	71	60 (16.4)	78	61.7 (19.7)	-+	-1.77[-7.57,4.03]
Penn 2009	33	59.9 (10.5)	32	52.2 (10.7)	-	7.7[2.54,12.86]
3.4.2 long term						
Durham 2003	19	93.5 (16.8)	21	87 (23.1)	+	6.5[-5.94,18.94]
Penn 2009	33	58.4 (11.2)	32	52.7 (10.1)		5.7[0.52,10.88]

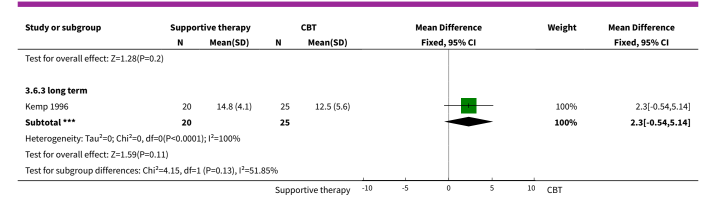
Analysis 3.5. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 5 Mental state: 3. Average endpoint general mental state score (BPRS, high = poor).

Study or subgroup	Suppor	tive therapy		СВТ	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
3.5.1 short term							
Haddock 1999	10	38.3 (17.4)	8	46.8 (8.8)	+	10.51%	-8.5[-20.87,3.87]
Kemp 1996	35	37.4 (8.5)	39	37.6 (10.1)		89.49%	-0.2[-4.44,4.04]
Subtotal ***	45		47			100%	-1.07[-5.08,2.94]
Heterogeneity: Tau ² =0; Chi ² =	1.55, df=1(P=0.2	1); I ² =35.37%					
Test for overall effect: Z=0.52	(P=0.6)						
3.5.2 medium term							
Pinto 1999	18	45.7 (11)	19	38.1 (9.7)		100%	7.6[0.9,14.3]
Subtotal ***	18		19			100%	7.6[0.9,14.3]
Heterogeneity: Tau ² =0; Chi ² =	0, df=0(P<0.0001	.); I ² =100%					
Test for overall effect: Z=2.22	(P=0.03)						
			Suppo	ortive therapy	-10 -5 0	5 10 CBT	

Analysis 3.6. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 6 Mental state: 4. Average endpoint general mental state score (BPRS short form, high = poor).

Study or subgroup	Suppor	tive therapy		СВТ	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
3.6.1 short term							
Kemp 1996	35	11.7 (3.3)	39	12.6 (5.8)		100%	-0.9[-3.02,1.22]
Subtotal ***	35		39		•	100%	-0.9[-3.02,1.22]
Heterogeneity: Not applicable							
Test for overall effect: Z=0.83(P=0.4	1)						
3.6.2 medium term							
Kemp 1996	31	16.7 (6.9)	36	14.5 (7.2)		100%	2.2[-1.18,5.58]
Subtotal ***	31		36			100%	2.2[-1.18,5.58]
Heterogeneity: Not applicable							
			Suppo	ortive therapy -10	-5 0 5	¹⁰ CBT	





Analysis 3.7. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 7 Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor).

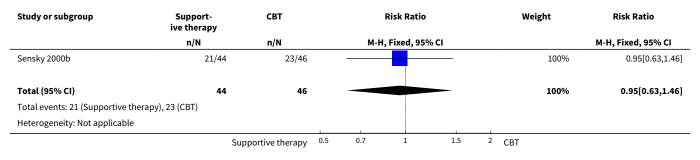
Mental state: 5. Average endpoint general mental state score (CPRS, skewed data, high = poor)

Study	Intervention	Intervention N		Mean	SD
		n	nedium term		
Sensky 2000b	Supportive therapy	44	22.9	17.3	
Sensky 2000b	CBT	46	20.5	13.1	
			long term		
Sensky 2000b	Supportive therapy	44	26.6	25.3	
Sensky 2000b	СВТ	46	15.1	12.0	

Analysis 3.8. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 8 Mental state: 6. Average endpoint general mental state score (PANSS general subscale, high = poor).

Study or subgroup	Supp	ortive therapy	СВТ		Mean Difference	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	Fixed, 95% CI
3.8.1 short term						
Levine 1998	6	31.8 (2.8)	6	14.7 (3.1)		17.1[13.76,20.44]
Penn 2009	33	30 (7.1)	32	25.6 (5.3)		4.4[1.36,7.44]
3.8.2 long term						
Penn 2009	33	29.6 (6.2)	32	26 (5.1)		3.6[0.84,6.36]
			S	upportive therapy	-20 -10 0 10 20	CBT

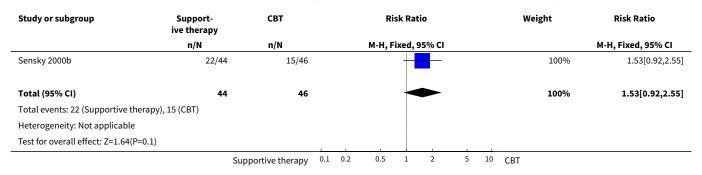
Analysis 3.9. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 9 Mental state: 7. No clinically important improvement in negative symptoms.



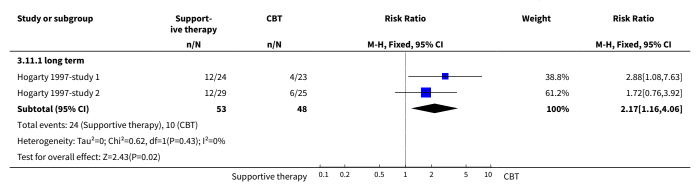


Study or subgroup	Support- CBT ive therapy				Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н	, Fixed, 95	% CI			M-H, Fixed, 95% CI
Test for overall effect: Z=0.22(P=0.83)			_			1			
		Supportive therapy	0.5	0.7	1	1.5	2	CBT	

Analysis 3.10. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 10 Mental state: 8. No clinically important improvement in depressive symptoms.



Analysis 3.11. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 11 Mental state: 9. Episode of affective symptoms.



Analysis 3.12. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 12 Mental state: 10. Average endpoint negative symptoms score (PANSS negative subscale, high = poor).

Study or subgroup Supportive		ortive therapy	erapy CBT			Mean Difference			ean Difference
N	N	Mean(SD)	N	Mean(SD)		Fixed, 95% CI			ixed, 95% CI
3.12.1 short term									
Levine 1998	6	15 (1.5)	6	7.8 (4)				>	7.2[3.78,10.62]
Penn 2009	33	12.7 (3.7)	32	12.4 (3.9)		-			0.3[-1.55,2.15]
3.12.2 long term									
Penn 2009	33	13.2 (4.9)	32	12.9 (4.4)					0.3[-1.96,2.56]
			S	Supportive therapy	-10	-5 0	5 1	.0 CBT	



Analysis 3.13. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 13 Mental state: 11. Average endpoint negative symptoms score (SANS, high = poor).

Study or subgroup	Suppor	tive therapy	СВТ			Mear	Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixe	ed, 95% CI		Fixed, 95% CI
3.13.1 medium term		·							
Pinto 1999	18	53.5 (19.1)	19	46.9 (19.4)				100%	6.6[-5.81,19.01]
Subtotal ***	18		19					100%	6.6[-5.81,19.01]
Heterogeneity: Not applicable									
Test for overall effect: Z=1.04(P=0.3)									
			Suppo	rtive therapy	-50	-25	0 25 50	CBT	

Analysis 3.14. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 14 Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor).

Mental state: 12. Average endpoint negative symptoms score (SANS, skewed data, high = poor)

	Melitat State. 12. Average	enuponit negative symptoms s	core (3AN3, 3Kewed data, mgm - p	001)
Study	Intervention	N	Mean	SD
		short term		
Tarrier 1998	Supportive therapy	21	10.19	5.83
Tarrier 1998	СВТ	23	9.83	4.58
		medium term		
Sensky 2000b	Supportive therapy	44	20.7	20.9
Sensky 2000b	CBT	46	22.0	17.0
		long term		
Sensky 2000b	Supportive therapy	44	25.1	27.5
Sensky 2000b	СВТ	46	18.2	17.7
Tarrier 1998	Supportive therapy	21	9.90	5.44
Tarrier 1998	СВТ	23	10.39	4.01

Analysis 3.15. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 15 Mental state: 13. Average endpoint positive symptoms score (PANSS positive subscale, high = poor).

Study or subgroup	Suppo	Supportive therapy CBT		СВТ	Mean Difference	Mean Difference	
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI	Fixed, 95% CI	
3.15.1 short term							
Levine 1998	6	13.7 (2.5)	6	7 (4.6)		6.7[2.51,10.89]	
Lewis 2002b	71	12.6 (4.8)	78	13 (5.1)		-0.45[-2.03,1.13]	
Penn 2009	33	16.5 (4)	32	14.2 (4)		2.3[0.35,4.25]	
3.15.2 long term							
Penn 2009	33	15.9 (3.6)	32	13.6 (3.4)		2.3[0.6,4]	
			S	upportive therapy	-10 -5 0 5 1	0 CBT	



Analysis 3.16. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 16 Mental state: 14. Average endpoint positive symptoms score (SAPS, skewed data, high = poor).

Mental state: 14. Average endpoint positive symptoms score (SAPS, skewed data, high = poor)

Study	Intervention Mean		n SD		N				
	medium term								
Pinto 1999	Supportive therapy	29.9	12.1	18					
Pinto 1999	CBT	17.9	17.0	19					

Analysis 3.17. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 17 Mental state: 15. Average endpoint positive symptoms score (BPRS, skewed data, high = poor).

Mental state: 15. Average endpoint positive symptoms score (BPRS, skewed data, high = poor)

Study	Intervention	N	Mean	SD						
short term										
Tarrier 1998	Supportive therapy	21	15.81	16.10						
Tarrier 1998	CBT	23	10.67	9.42						
		long term								
Tarrier 1998	Supportive therapy	21	16.30	16.71	•					
Tarrier 1998	CBT	23	10.59	11.10						

Analysis 3.18. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 18 Mental state: 16. Average endpoint score (PANSS thought disturbance cluster, high = poor).

Study or subgroup	Supportive therapy		CBT			Mean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed, 95% CI			Fixed, 95% CI
3.18.1 short term									
Levine 1998	6	7.8 (2.1)	6	3.5 (3.3)			-	100%	4.3[1.17,7.43]
Subtotal ***	6		6					100%	4.3[1.17,7.43]
Heterogeneity: Not applicable									
Test for overall effect: Z=2.69(P=0.01)								
			Suppo	ortive therapy	-10	-5	0 5	¹⁰ CBT	

Analysis 3.19. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 19 Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor).

Mental state: 17. Average endpoint delusions score (PSYRATS delusions score, skewed data, high = poor)

Study	Intervention	Mean	SD	N
		short term		
Lewis 2002b	Supportive therapy	6.13	6.98	67
Lewis 2002b	CBT	6.95	7.66	74
Penn 2009	Supportive therapy	10.4	5.9	33
Penn 2009	СВТ	8	7.7	32
		long term		
Durham 2003	Supportive therapy	9.7	6.1	19
Durham 2003	СВТ	11.1	5.8	21
Penn 2009	Supportive therapy	9.0	6.8	33
Penn 2009	СВТ	s6.9	7.0	32

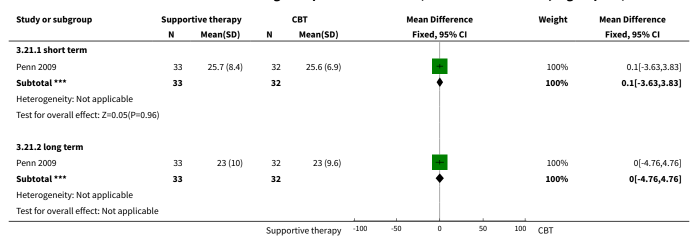


Analysis 3.20. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 20 Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor).

Mental state: 18. Average endpoint hallucinations score (PSYRATS hallucinations score, skewed data, high = poor)

Study	Intervention	Mean	SD	N	
		long term			
Durham 2003	Supportive therapy	18.0	12.2	19	
Durham 2003	CBT	18.5	12.8	20	

Analysis 3.21. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 21 Mental state: 19. Average endpoint voices score (PSYRATS voices score, high = poor).



Analysis 3.22. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 22 Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data).

Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data)

Study	Follow-up length	Intervention	Mean	SD	N
		Mal	evolence		
Penn 2009	Short term	Supportive therapy	8.1	5.4	33
Penn 2009	Short term	CBT	6.3	5.4	32
Penn 2009	Long term	Supportive therapy	6.7	4.9	33
Penn 2009	Long term	CBT	6.3	5.4	32
		Ben	evolence		
Penn 2009	Short term	Supportive therapy	5.7	5.5	33
Penn 2009	Short term	CBT	4.4	5.3	32
Penn 2009	Long term	Supportive therapy	6.2	5.4	33
Penn 2009	Long term	CBT	5.3	5.2	32
		Re	sistance		
Penn 2009	Short term	Supportive therapy	16.3	6.6	33
Penn 2009	Short term	CBT	15	7.3	32
Penn 2009	Long term	Supportive therapy	15.3	7.3	33
Penn 2009	Long term	CBT	13.9	7.8	32
		Eng	agement		
Penn 2009	Short term	Supportive therapy	7.8	7.4	33
Penn 2009	Short term	CBT	5.4	5.7	32
Penn 2009	Long term	Supportive therapy	7.3	6.8	33
Penn 2009	Long term	CBT	6.4	7.2	32
		Omi	nipotence		



Mental state: 20. Average endpoint beliefs about voices score (BAVQ, high = poor, skewed data)

Study	Follow-up length	Intervention	Mean	SD	N
Penn 2009	Short term	Supportive therapy	8.6	5	33
Penn 2009	Short term	CBT	8	4.2	32
Penn 2009	Long term	Supportive therapy	7.7	4.3	33
Penn 2009	Long term	CBT	7.3	3.3	32

Analysis 3.23. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 23 Mental state: 21. Average endpoint depression score (MADRS, skewed data, high = poor).

Mental state: 21. Average endpoint depression score (MADRS, skewed data, high = poor)

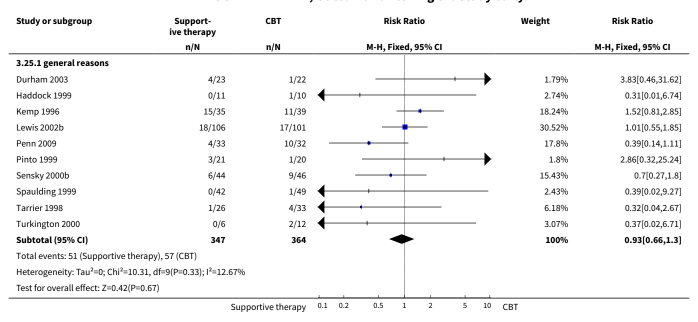
Study	Intervention	N	Mean	SD	
		medium ter	m		
Sensky 2000b	Supportive therapy	44	6.0	5.4	
Sensky 2000b	CBT	46	4.8	3.5	
	'	long term			
Sensky 2000b	Supportive therapy	44	6.7	7.1	
Sensky 2000b	CBT	46	3.7	3.2	•

Analysis 3.24. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 24 Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor).

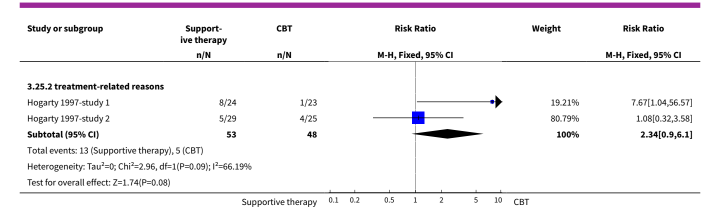
Mental state: 22. Average endpoint depression score (BDI-II, skewed data, high = poor)

Study	Follow-up length	Intervention	Mean	SD	N
Penn 2009	Short term	Supportive therapy	13.9	10.7	33
Penn 2009	Short term	CBT	10.5	8.5	32
Penn 2009	Long term	Supportive therapy	17.9	13.6	33
Penn 2009	Long term	CBT	11.5	9.4	32

Analysis 3.25. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 25 Leaving the study early.







Analysis 3.26. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 26 General functioning: 1. Average endpoint general functioning score (GAF & GAS, high = good).

Study or subgroup	Supp	ortive therapy		CBT	Mean Difference	Mean Difference
	N	N Mean(SD)		Mean(SD)	Fixed, 95% CI	Fixed, 95% CI
3.26.1 short term						
Kemp 1996	33	44.5 (10.4)	37	54 (17.3)		-9.5[-16.11,-2.89]
3.26.2 medium term						
Kemp 1996	31	43.3 (10.6)	36	55.9 (17.5)		-12.6[-19.43,-5.77]
3.26.3 long term						
Durham 2003	12	36.3 (9.8)	18	35.8 (9.7)		0.5[-6.63,7.63]
Kemp 1996	23	48.3 (14.5)	25	62.8 (18.4)		-14.5[-23.83,-5.17]
				CBT	-20 -10 0 10 20	Supportive therapy

Analysis 3.27. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 27 General functioning: 2. Average endpoint social functioning score (SFS, high = good).

Study or subgroup	Suppo	rtive therapy		СВТ		Mean Difference	•	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fixed, 95% CI			Fixed, 95% CI
3.27.1 short term									
Penn 2009	33	121.9 (23.3)	32	129.1 (20.5)		-		100%	-7.2[-17.86,3.46]
Subtotal ***	33		32			•		100%	-7.2[-17.86,3.46]
Heterogeneity: Not applicable									
Test for overall effect: Z=1.32(P=0.	19)								
3.27.2 long term									
Penn 2009	33	119.7 (24.2)	32	128.5 (28.5)		-		100%	-8.8[-21.67,4.07]
Subtotal ***	33		32			•		100%	-8.8[-21.67,4.07]
Heterogeneity: Not applicable									
Test for overall effect: Z=1.34(P=0.	18)								
Test for subgroup differences: Chi	² =0.04, df=	1 (P=0.85), I ² =0%			1				
				CBT	-100 -5	50 0	50 100	Supportive	therapy



Analysis 3.28. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 28 Satisfaction with treatment: Recipient of care not satisfied with treatment.

Study or subgroup	Support- ive therapy	СВТ			Risk Ratio	•		Weig	ht	Risk Ratio
	n/N	n/N		М-Н	, Fixed, 95	% CI				M-H, Fixed, 95% CI
3.28.1 long term										
Durham 2003	10/23	3/22				-			100%	3.19[1.01,10.07]
Subtotal (95% CI)	23	22				-			100%	3.19[1.01,10.07]
Total events: 10 (Supportive th	nerapy), 3 (CBT)									
Heterogeneity: Not applicable										
Test for overall effect: Z=1.98(F	P=0.05)									
	Sup	portive therapy	0.01	0.1	1	10	100	CBT		

Analysis 3.29. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 29 Quality of life: Average endpoint score (RSES, high = good).

Study or subgroup	Suppor	tive therapy		CBT	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
3.29.1 short term							
Penn 2009	33	28.6 (6.2)	32	29.4 (6)	+	100%	-0.8[-3.77,2.17]
Subtotal ***	33		32		▼	100%	-0.8[-3.77,2.17]
Heterogeneity: Not applicable	e						
Test for overall effect: Z=0.53((P=0.6)						
3.29.2 long term							
Penn 2009	33	27.6 (6.7)	32	29.3 (7.6)	+	100%	-1.7[-5.19,1.79]
Subtotal ***	33		32		♦	100%	-1.7[-5.19,1.79]
Heterogeneity: Tau ² =0; Chi ² =0	0, df=0(P<0.0001	.); I ² =100%					
Test for overall effect: Z=0.96	(P=0.34)						
Test for subgroup differences	:: Chi²=0.15, df=1	(P=0.7), I ² =0%					
	·			CBT	-50 -25 0 25 50	Supportive	therapy

Analysis 3.30. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 30 Death.

1/26 26	n/N 1/33 33		M-H, Fixe	d, 95% CI		100% 100%	M-H, Fixed, 95% CI
	•						
	•						
26	33					1000/	
						100%	1.27[0.08,19.34]
1/23	0/22			-	-	100%	2.88[0.12,67.03]
23	22				-	100%	2.88[0.12,67.03]
	23	23 22	23 22	23 22	23 22	23 22	23 22 100%



Study or subgroup	Support- ive therapy	СВТ	Risk Ratio M-H, Fixed, 95% CI			Weight	Risk Ratio		
	n/N	n/N					M-H, Fixed, 95% CI		
Test for overall effect: Z=0.66(P=0.51)						1			
		Supportive therapy	0.002	0.1	1	10	500	CBT	

Analysis 3.31. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 31 Insight:1. Average endpoint insight score (SAI, skewed data, high = good).

Insight:1. Average endpoint insight score (SAI, skewed data, high = good)

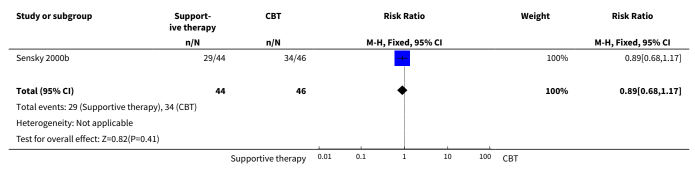
Study	Intervention	Mean	SD	N							
short term											
Kemp 1996	Supportive therapy	40.60	31.20	35							
Kemp 1996	CBT	63.00	23.60	39							
medium term											
Kemp 1996	Supportive therapy	41.90	30.80	29							
Kemp 1996	CBT	62.60	23.50	34							
		long tern	n								
Kemp 1996	Supportive therapy	42.60	36.50	20							
Kemp 1996	CBT	63.40	25.50	30							
				'							

Analysis 3.32. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 32 Insight: 2. Average endpoint insight score (BCIS composite, skewed data, high = good).

Insight: 2. Average endpoint insight score (BCIS composite, skewed data, high = good)

Study	Follow-up length	Intervention	Mean	SD	N
Penn 2009	Short term	Supportive therapy	4.7	5.4	33
Penn 2009	Short term	CBT	4.2	6.6	32
Penn 2009	Long term	Supportive therapy	3.7	4.4	33
Penn 2009	Long term	CBT	4.4	5.6	32

Analysis 3.33. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 33 Medication: no reduction in dose of antipsychotic medication.





Analysis 3.34. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 34 Attitude to medication: 1. Average endpoint attitude to medication score (AMQ, high = good).

Study or subgroup	Supportive therapy		CBT			Mea	n Differen	:e		Weight	Mean Difference	
	N	Mean(SD)	N Mean(SD)		Fixed, 95% CI						Fixed, 95% CI	
3.34.1 short term												
Kemp 1996	35	14.9 (6.1)	39	19.4 (3.7)						100%	-4.5[-6.83,-2.17]	
Subtotal ***	35		39							100%	-4.5[-6.83,-2.17]	
Heterogeneity: Not applicable												
Test for overall effect: Z=3.78(P=0)												
				CBT	-10	-5	0	5	10	Supportive t	:herapy	

Analysis 3.35. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 35 Attitude to medication: 2. Average endpoint attitude to medication score (DAI, high = good).

Study or subgroup	Suppor	tive therapy		СВТ	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI		Fixed, 95% CI
3.35.1 short term							
Kemp 1996	28	44.4 (8.1)	35	50.1 (6.3)		100%	-5.7[-9.35,-2.05]
Subtotal ***	28		35			100%	-5.7[-9.35,-2.05]
Heterogeneity: Not applicable							
Test for overall effect: Z=3.06(P=0)							
3.35.2 long term							
Kemp 1996	16	44.6 (7.5)	28	49.5 (6.9)		100%	-4.9[-9.38,-0.42]
Subtotal ***	16		28			100%	-4.9[-9.38,-0.42]
Heterogeneity: Not applicable							
Test for overall effect: Z=2.15(P=0.0	3)						
Test for subgroup differences: Chi ² =	0.07, df=1	(P=0.79), I ² =0%					
				CBT	-10 -5 0 5	10 Supportive	therapy

Analysis 3.36. Comparison 3 SUPPORTIVE THERAPY versus COGNITIVE BEHAVIOURAL THERAPY, Outcome 36 Economic outcomes: Direct costs (skewed data).

Economic outcomes: Direct costs (skewed data)

Study	Intervention	Mea	ın	SD	N						
medium term											
Kemp 1996	Supportive therapy	252	234	34							
Kemp 1996	СВТ	187	292	36							
		long to	erm								
Kemp 1996	Supportive therapy	326	404	21							
Kemp 1996	СВТ	239	281	24							

Comparison 4. SUPPORTIVE THERAPY versus FAMILY THERAPY

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: 1. Relapse	2		Risk Ratio (M-H, Fixed, 95% CI)	Totals not selected



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.1 long term	2		Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
2 Global state: 2. No remission	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.87 [1.11, 3.15]
3 Service outcomes: Hospitalisation	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.93 [0.89, 4.17]
4 Mental state: Episode of af- fective symptoms	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.1 long term	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.71 [0.82, 3.60]
5 Leaving the study early (by long term)	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
5.1 general reasons	1	39	Risk Ratio (M-H, Fixed, 95% CI)	0.70 [0.13, 3.75]
5.2 treatment-related reasons	1	48	Risk Ratio (M-H, Fixed, 95% CI)	1.6 [0.61, 4.19]
6 General functioning: 1. No paid work	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
6.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.57, 1.63]
7 General functioning: 2. Admission to residential placement	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
7.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.24, 4.59]
8 General fuctioning: 3. Admission to jail	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
8.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.24, 4.59]
9 Behaviour: 1. Social impairment on SBAS	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
9.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.46 [1.04, 2.04]
10 Behaviour: 2. Poor coping style with relatives	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
10.1 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	0.90 [0.37, 2.20]
11 Engagement with services: Poor attendance at appoint- ments	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.93 [0.89, 4.17]
12 Compliance: Poor compliance to therapy	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only

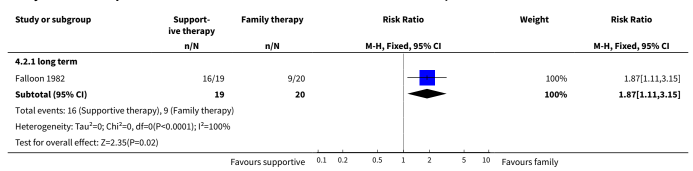


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
12.1 medium term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	2.63 [1.30, 5.35]
12.2 long term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	1.29 [0.69, 2.39]
13 Medication: Prescribed IM depot medication	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
13.1 medium term	1	39	Risk Ratio (M-H, Fixed, 95% CI)	2.11 [0.99, 4.47]

Analysis 4.1. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 1 Global state: 1. Relapse.

Study or subgroup	Supportive therapy	Family therapy	Risk Ratio	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
4.1.1 long term				
Falloon 1982	16/19	5/20		3.37[1.54,7.38]
Hogarty 1997-study 1	15/24	15/24	· · · · · · · · · · · · · · · · · · ·	1[0.65,1.55]
		Favours supportive 0.1	0.2 0.5 1 2 5	10 Favours family

Analysis 4.2. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 2 Global state: 2. No remission.

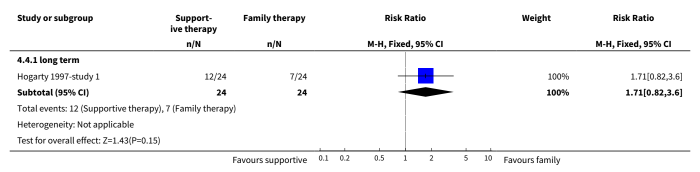


Analysis 4.3. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 3 Service outcomes: Hospitalisation.

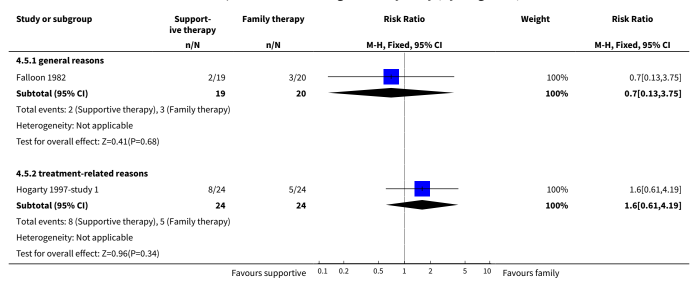
Study or subgroup	Support- ive therapy	Family therapy		Risk Ratio			Weight	Risk Ratio			
	n/N	n/N			M-H, F	ixed,	95% CI				M-H, Fixed, 95% CI
4.3.1 long term											
Falloon 1982	11/19	6/20				+	-	_		100%	1.93[0.89,4.17]
Subtotal (95% CI)	19	20				-		-		100%	1.93[0.89,4.17]
Total events: 11 (Supportive therapy	y), 6 (Family therapy)									
Heterogeneity: Not applicable											
Test for overall effect: Z=1.67(P=0.09)										
	1	Favours supportive	0.1	0.2	0.5	1	2	5	10	Favours family	



Analysis 4.4. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 4 Mental state: Episode of affective symptoms.



Analysis 4.5. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 5 Leaving the study early (by long term).

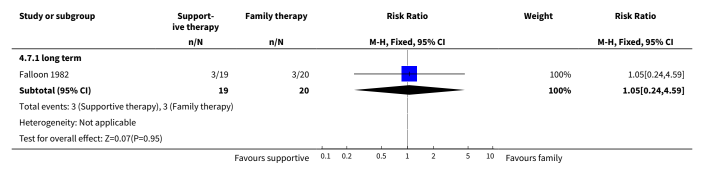


Analysis 4.6. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 6 General functioning: 1. No paid work.

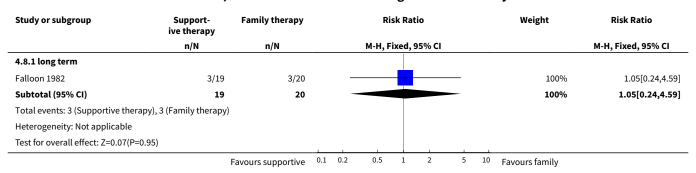
Study or subgroup	Support- ive therapy	Family therapy			Ri	sk Rati	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	5% CI				M-H, Fixed, 95% CI
4.6.1 long term											
Falloon 1982	11/19	12/20			_		_			100%	0.96[0.57,1.63]
Subtotal (95% CI)	19	20			4	~	-			100%	0.96[0.57,1.63]
Total events: 11 (Supportive therap	y), 12 (Family therap	y)									
Heterogeneity: Not applicable											
Test for overall effect: Z=0.13(P=0.89	9)										
	1	Favours supportive	0.1	0.2	0.5	1	2	5	10	Favours family	



Analysis 4.7. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 7 General functioning: 2. Admission to residential placement.



Analysis 4.8. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 8 General fuctioning: 3. Admission to jail.

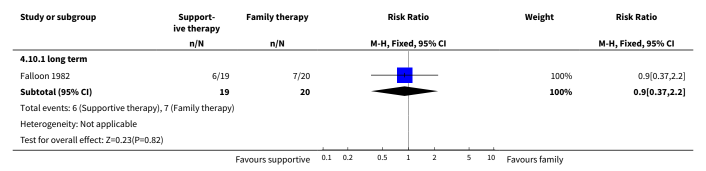


Analysis 4.9. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 9 Behaviour: 1. Social impairment on SBAS.

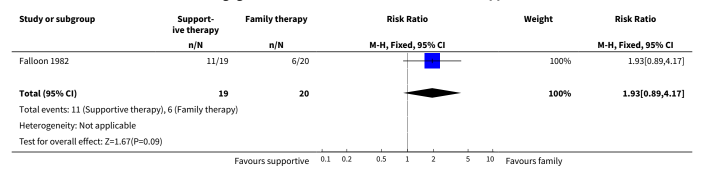
Study or subgroup	Support- ive therapy				Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	l, Fixed, 95%	CI			M-H, Fixed, 95% CI
4.9.1 long term									
Falloon 1982	18/19	13/20				-	\rightarrow	100%	1.46[1.04,2.04]
Subtotal (95% CI)	19	20			-	-		100%	1.46[1.04,2.04]
Total events: 18 (Supportive th	erapy), 13 (Family therap	y)							
Heterogeneity: Not applicable									
Test for overall effect: Z=2.18(P	2=0.03)								
		Favours supportive	0.5	0.7	1	1.5	2	Favours family	



Analysis 4.10. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 10 Behaviour: 2. Poor coping style with relatives.



Analysis 4.11. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 11 Engagement with services: Poor attendance at appointments.

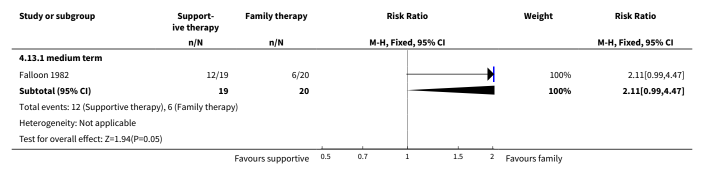


Analysis 4.12. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 12 Compliance: Poor compliance to therapy.

Study or subgroup	Support- ive therapy	Family therapy n/N		/ Risk Ratio					Weight	Risk Ratio
	n/N			M-	H, Fixed	, 95% CI				M-H, Fixed, 95% CI
4.12.1 medium term										
Falloon 1982	15/19	6/20				-			100%	2.63[1.3,5.35
Subtotal (95% CI)	19	20					-		100%	2.63[1.3,5.35
Total events: 15 (Supportive therapy)), 6 (Family therapy	·)								
Heterogeneity: Not applicable										
Test for overall effect: Z=2.68(P=0.01))									
4.12.2 long term										
Falloon 1982	11/19	9/20			_	-			100%	1.29[0.69,2.39
Subtotal (95% CI)	19	20			4	>			100%	1.29[0.69,2.39
Total events: 11 (Supportive therapy)), 9 (Family therapy	·)								
Heterogeneity: Not applicable										
Test for overall effect: Z=0.8(P=0.42)										
		Favours supportive	0.1	0.2 0	.5 1	2	5	¹⁰ Favo	urs family	



Analysis 4.13. Comparison 4 SUPPORTIVE THERAPY versus FAMILY THERAPY, Outcome 13 Medication: Prescribed IM depot medication.



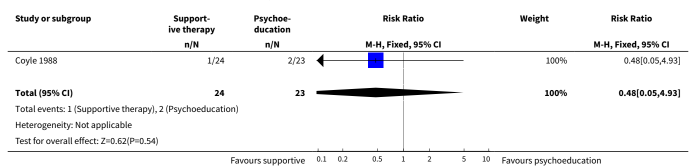
Comparison 5. SUPPORTIVE THERAPY versus PSYCHOEDUCATION

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Service outcomes: Hospitalisation	1	47	Risk Ratio (M-H, Fixed, 95% CI)	0.48 [0.05, 4.93]
2 Mental state. 1. No clinically important improvement in general mental state	1	19	Risk Ratio (M-H, Fixed, 95% CI)	1.61 [0.96, 2.68]
3 Mental state: 2. Average endpoint general score (PANSS general subscale, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	2.86 [-3.21, 8.93]
4 Mental state: 3. Average endpoint negative symptoms score (PANSS negative subscale, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	1.70 [-4.00, 7.40]
5 Mental state: 4. Average endpoint positive symptoms score (PANSS positive subscale, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	2.71 [-0.71, 6.13]
6 Mental state: 5. Average endpoint de- pression score (CDRS, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	1.47 [-1.35, 4.29]
7 Leaving the study early	2	71	Risk Ratio (M-H, Fixed, 95% CI)	0.57 [0.21, 1.54]
8 Quality of life: Average endpoint quality of life score (QLS, high = good)	1	19	Mean Difference (IV, Fixed, 95% CI)	-0.07 [-21.11, 20.97]
9 Death	1	47	Risk Ratio (M-H, Fixed, 95% CI)	2.88 [0.12, 67.29]
10 Behaviour: Average endpoint score (composed of ROMI and ITAQ items, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
10.1 Need for treatment	1	19	Mean Difference (IV, Fixed, 95% CI)	-0.02 [-0.44, 0.40]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
10.2 Benefits of medication	1	19	Mean Difference (IV, Fixed, 95% CI)	-0.19 [-0.56, 0.18]
11 Insight: Average endpoint treatment attitude score (ITAQ, high = good)	1	19	Mean Difference (IV, Fixed, 95% CI)	-1.55 [-5.85, 2.75]
12 Compliance: 1. Poor compliance to therapy	1	19	Odds Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
13 Compliance: 2. Adherence to medication (self-report)	1	19	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.83, 1.21]
14 Compliance: 3. Average endpoint adherence score (ROMI, high = good)	1	19	Mean Difference (IV, Fixed, 95% CI)	0.45 [-3.58, 4.48]
15 Compliance: 4. Average endpoint non- adherence score (ROMI, high = poor)	1	19	Mean Difference (IV, Fixed, 95% CI)	1.75 [-0.61, 4.11]

Analysis 5.1. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 1 Service outcomes: Hospitalisation.



Analysis 5.2. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 2 Mental state. 1. No clinically important improvement in general mental state.

Study or subgroup	Support- ive therapy	Psychoe- ducation			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H	l, Fixed, 95% C	I			M-H, Fixed, 95% CI
Uzenoff 2007	9/9	6/10			-			100%	1.61[0.96,2.68]
Total (95% CI)	9	10			•			100%	1.61[0.96,2.68]
Total events: 9 (Supportive therapy)	, 6 (Psychoeducation)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.82(P=0.07	7)								
	Fav	ours supportive	0.01	0.1	1	10	100	Favours psychoeducation	on



Analysis 5.3. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 3 Mental state: 2. Average endpoint general score (PANSS general subscale, high = poor).

Study or subgroup	Suppor	tive therapy	Psych	oeducation	Mean Difference					Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% CI				Fixed, 95% CI	
Uzenoff 2007	9	27.6 (8.6)	10	24.7 (3.7)			+			100%	2.86[-3.21,8.93]	
Total ***	9		10				•			100%	2.86[-3.21,8.93]	
Heterogeneity: Not applicable												
Test for overall effect: Z=0.92(P=0.36)											
			Favou	irs supportive	-100	-50	0	50	100	Favours psy	/choeducation	

Analysis 5.4. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 4 Mental state: 3. Average endpoint negative symptoms score (PANSS negative subscale, high = poor).

Study or subgroup	udy or subgroup Supportive thera			oeducation		Ме	an Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C				Fixed, 95% CI
Uzenoff 2007	9	16 (7.4)	10	14.3 (4.9)			+			100%	1.7[-4,7.4]
Total ***	9		10				•			100%	1.7[-4,7.4]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.58(P=0.56	6)										
			Favou	rs supportive	-100	-50	0	50	100	Favours psy	choeducation

Analysis 5.5. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 5 Mental state: 4. Average endpoint positive symptoms score (PANSS positive subscale, high = poor).

Study or subgroup	Supportive therapy		Psych	oeducation		Me	an Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		Fi	xed, 95% C	I			Fixed, 95% CI
Uzenoff 2007	9	12.1 (4.6)	10	9.4 (2.7)			+			100%	2.71[-0.71,6.13]
Total ***	9		10				♦			100%	2.71[-0.71,6.13]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.55(P=0.12	2)				1						
			Favou	rs supportive	-100	-50	0	50	100	Favours psy	choeducation

Analysis 5.6. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 6 Mental state: 5. Average endpoint depression score (CDRS, high = poor).

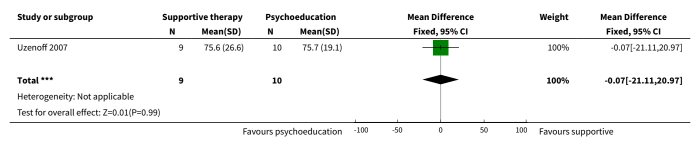
Study or subgroup	Suppor	tive therapy	Psych	oeducation	Mean Difference					Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% CI				Fixed, 95% CI	
Uzenoff 2007	9	11.7 (3.9)	10	10.2 (1.9)			+			100%	1.47[-1.35,4.29]	
Total ***	9		10				•			100%	1.47[-1.35,4.29]	
Heterogeneity: Not applicable												
Test for overall effect: Z=1.02(P=0.32	L)											
			Favou	rs supportive	-100	-50	0	50	100	Favours psy	choeducation	



Analysis 5.7. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 7 Leaving the study early.

Study or subgroup	Support- ive therapy	Psychoe- ducation			Ri	sk Rat	io		Weight	Risk Ratio	
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
Coyle 1988	3/24	6/23	_		-		_			69.02%	0.48[0.14,1.69]
Uzenoff 2007	2/11	3/13				•		-		30.98%	0.79[0.16,3.9]
Total (95% CI)	35	36		-			_			100%	0.57[0.21,1.54]
Total events: 5 (Supportive th	erapy), 9 (Psychoeducation)										
Heterogeneity: Tau ² =0; Chi ² =0	0.23, df=1(P=0.63); I ² =0%										
Test for overall effect: Z=1.1(P	=0.27)										
	Fav	ours supportive	0.1	0.2	0.5	1	2	5	10	Favours psychoeducat	ion

Analysis 5.8. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 8 Quality of life: Average endpoint quality of life score (QLS, high = good).



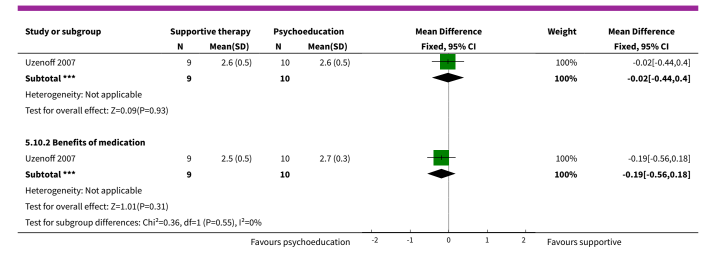
Analysis 5.9. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 9 Death.

Study or subgroup	Support- ive therapy	Psychoe- ducation			Ris	k Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, Fi	xed, 9	5% CI			I	M-H, Fixed, 95% CI
Coyle 1988	1/24	0/23	_				1		→	100%	2.88[0.12,67.29]
Total (95% CI)	24	23	_							100%	2.88[0.12,67.29]
Total events: 1 (Supportive therapy),	0 (Psychoeducation)										
Heterogeneity: Not applicable											
Test for overall effect: Z=0.66(P=0.51)											
	Favo	ours supportive	0.1	0.2	0.5	1	2	5	10	Favours psychoeducation	n

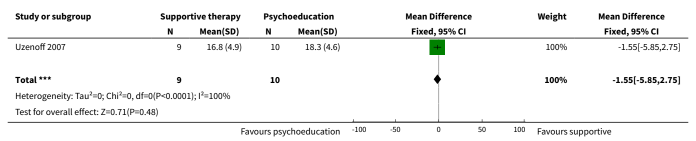
Analysis 5.10. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 10 Behaviour: Average endpoint score (composed of ROMI and ITAQ items, high = good).

Study or subgroup	Suppo	pportive therapy Psychoeducatio		oeducation		Меа	an Differe	nce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Fixed, 95% CI				Fixed, 95% CI		
5.10.1 Need for treatment											
		Fa	vours psychoeducation		-2	-1	0	1	2	Favours supp	ortive





Analysis 5.11. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 11 Insight: Average endpoint treatment attitude score (ITAQ, high = good).



Analysis 5.12. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 12 Compliance: 1. Poor compliance to therapy.

Study or subgroup	Support- ive therapy	Psychoe- ducation		Odds Ratio		Weight		Odds Ratio	
	n/N	n/N		М-Н	I, Fixed, 95%	CI			M-H, Fixed, 95% CI
Uzenoff 2007	0/9	0/10							Not estimable
Total (95% CI)	9	10							Not estimable
Total events: 0 (Supportive the	erapy), 0 (Psychoeducation)								
Heterogeneity: Not applicable									
Test for overall effect: Not app	licable								
	Fa	vours supportive	0.01	0.1	1	10	100	Favours psychoeducation	on



Analysis 5.13. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 13 Compliance: 2. Adherence to medication (self-report).

Study or subgroup	up Support- Psychoe- Risk Ratio ive therapy ducation			Weight	Risk Ratio				
	n/N	n/N		М-Н	, Fixed, 95% (CI			M-H, Fixed, 95% CI
Uzenoff 2007	9/9	10/10			+			100%	1[0.83,1.21]
Total (95% CI)	9	10			•			100%	1[0.83,1.21]
Total events: 9 (Supportive therapy),	, 10 (Psychoeducation)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable									
	Fav	ours supportive	0.01	0.1	1	10	100	Favours psychoeducation	on

Analysis 5.14. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 14 Compliance: 3. Average endpoint adherence score (ROMI, high = good).

Study or subgroup	Suppor	tive therapy	Psych	oeducation		Ме	an Differen	ce		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% C	:1			Fixed, 95% CI
Uzenoff 2007	9	16.8 (3.7)	10	16.3 (5.2)			+			100%	0.45[-3.58,4.48]
Total ***	9		10				•			100%	0.45[-3.58,4.48]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.22(P=0.83)										
		Fav	ours psy	choeducation	-100	-50	0	50	100	Favours supp	ortive

Analysis 5.15. Comparison 5 SUPPORTIVE THERAPY versus PSYCHOEDUCATION, Outcome 15 Compliance: 4. Average endpoint nonadherence score (ROMI, high = poor).

Study or subgroup	Suppor	tive therapy	Psych	oeducation	Mean Difference		Weight		Mean Difference		
	N	Mean(SD)	N	Mean(SD)		Fi	ixed, 95% CI				Fixed, 95% CI
Uzenoff 2007	9	12.1 (2.5)	10	10.4 (2.8)			+			100%	1.75[-0.61,4.11]
Total ***	9		10				•			100%	1.75[-0.61,4.11]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.45(P=0.1	5)										
		Fav	ours psy	choeducation	-100	-50	0	50	100	Favours suppor	tive

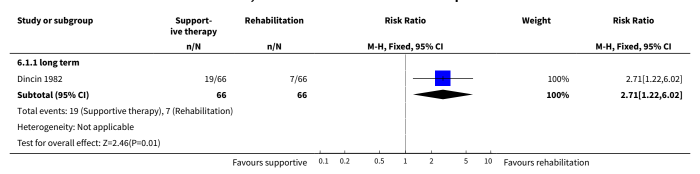
Comparison 6. SUPPORTIVE THERAPY versus REHABILITATION PROGRAMME

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Service outcomes: Hospitalisation	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 long term	1	132	Risk Ratio (M-H, Fixed, 95% CI)	2.71 [1.22, 6.02]

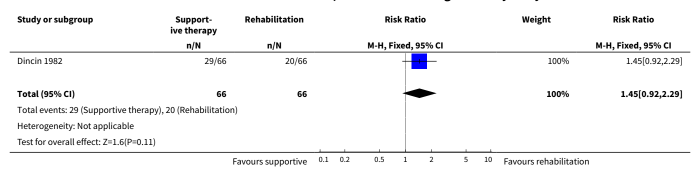


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2 Leaving the study early	1	132	Risk Ratio (M-H, Fixed, 95% CI)	1.45 [0.92, 2.29]
3 General functioning: No gainful employment	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 long term	1	132	Risk Ratio (M-H, Fixed, 95% CI)	1.04 [0.85, 1.29]

Analysis 6.1. Comparison 6 SUPPORTIVE THERAPY versus REHABILITATION PROGRAMME, Outcome 1 Service outcomes: Hospitalisation.



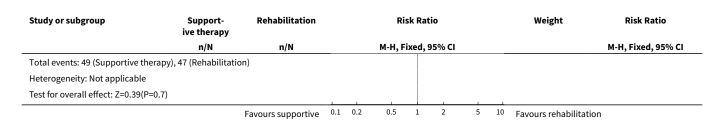
Analysis 6.2. Comparison 6 SUPPORTIVE THERAPY versus REHABILITATION PROGRAMME, Outcome 2 Leaving the study early.



Analysis 6.3. Comparison 6 SUPPORTIVE THERAPY versus REHABILITATION PROGRAMME, Outcome 3 General functioning: No gainful employment.

Study or subgroup	Support- ive therapy	Rehabilitation			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
6.3.1 long term											
Dincin 1982	49/66	47/66				-				100%	1.04[0.85,1.29]
Subtotal (95% CI)	66	66				*				100%	1.04[0.85,1.29]
		Favours supportive	0.1	0.2	0.5	1	2	5	10	Favours rehabilitation	

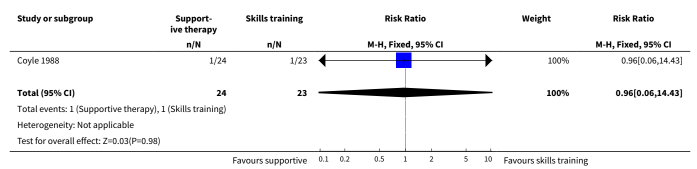




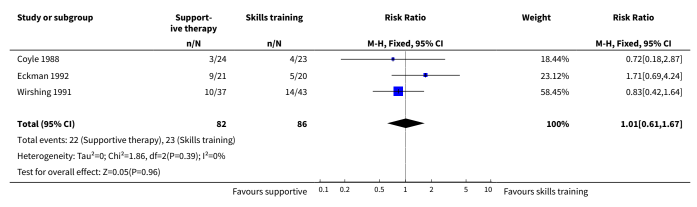
Comparison 7. SUPPORTIVE THERAPY versus SKILLS TRAINING

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Service outcomes: Hospitalisation	1	47	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.06, 14.43]
2 Leaving the study early	3	168	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.61, 1.67]
3 Death	1	47	Risk Ratio (M-H, Fixed, 95% CI)	2.88 [0.12, 67.29]

Analysis 7.1. Comparison 7 SUPPORTIVE THERAPY versus SKILLS TRAINING, Outcome 1 Service outcomes: Hospitalisation.



Analysis 7.2. Comparison 7 SUPPORTIVE THERAPY versus SKILLS TRAINING, Outcome 2 Leaving the study early.





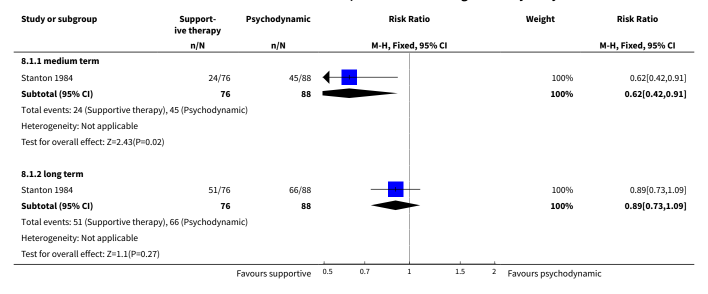
Analysis 7.3. Comparison 7 SUPPORTIVE THERAPY versus SKILLS TRAINING, Outcome 3 Death.

Study or subgroup	Support- ive therapy	Skills training			Ri	sk Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
Coyle 1988	1/24	0/23	_				1		→	100%	2.88[0.12,67.29]
Total (95% CI)	24	23	-							100%	2.88[0.12,67.29]
Total events: 1 (Supportive therapy	r), 0 (Skills training)										
Heterogeneity: Not applicable											
Test for overall effect: Z=0.66(P=0.5	1)										
	F	avours supportive	0.1	0.2	0.5	1	2	5	10	Favours skills training	

Comparison 8. SUPPORTIVE THERAPY versus PSYCHODYNAMIC PSYCHOTHERAPY

Outcome or subgroup ti- tle	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Leaving the study early	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 medium term	1	164	Risk Ratio (M-H, Fixed, 95% CI)	0.62 [0.42, 0.91]
1.2 long term	1	164	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.73, 1.09]

Analysis 8.1. Comparison 8 SUPPORTIVE THERAPY versus PSYCHODYNAMIC PSYCHOTHERAPY, Outcome 1 Leaving the study early.

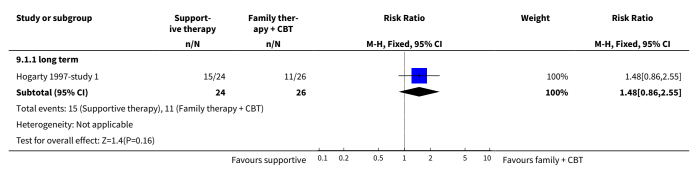




Comparison 9. SUPPORTIVE THERAPY versus COMBINATION OF OTHER PSYCHOSOCIAL INTERVENTIONS

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: Relapse	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 long term	1	50	Risk Ratio (M-H, Fixed, 95% CI)	1.48 [0.86, 2.55]
2 Mental state: Episode of affective symptoms	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 long term	1	50	Risk Ratio (M-H, Fixed, 95% CI)	1.63 [0.81, 3.28]
3 Leaving the study early: Treat- ment-related reasons	1	50	Risk Ratio (M-H, Fixed, 95% CI)	8.67 [1.17, 64.26]

Analysis 9.1. Comparison 9 SUPPORTIVE THERAPY versus COMBINATION OF OTHER PSYCHOSOCIAL INTERVENTIONS, Outcome 1 Global state: Relapse.

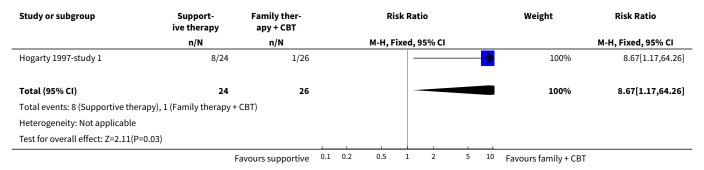


Analysis 9.2. Comparison 9 SUPPORTIVE THERAPY versus COMBINATION OF OTHER PSYCHOSOCIAL INTERVENTIONS, Outcome 2 Mental state: Episode of affective symptoms.

Study or subgroup	Support- ive therapy	Family ther- apy + CBT			Ri	sk Rat	io:			Weight	Risk Ratio
	n/N	n/N			M-H, F	ixed, 9	95% CI				M-H, Fixed, 95% CI
9.2.1 long term											
Hogarty 1997-study 1	12/24	8/26				+	+			100%	1.63[0.81,3.28]
Subtotal (95% CI)	24	26				-	-			100%	1.63[0.81,3.28]
Total events: 12 (Supportive th	herapy), 8 (Family therapy +	- CBT)									
Heterogeneity: Tau ² =0; Chi ² =0	, df=0(P<0.0001); I ² =100%										
Test for overall effect: Z=1.36(F	P=0.18)										
	Fi	avours supportive	0.1	0.2	0.5	1	2	5	10	Favours family + CBT	



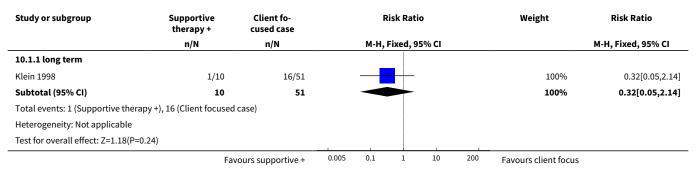
Analysis 9.3. Comparison 9 SUPPORTIVE THERAPY versus COMBINATION OF OTHER PSYCHOSOCIAL INTERVENTIONS, Outcome 3 Leaving the study early: Treatment-related reasons.



Comparison 10. SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus CLIENT-FOCUSED CASE MANAGEMENT

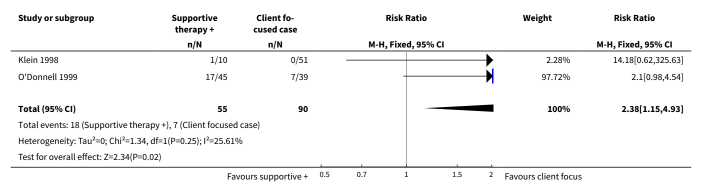
Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: Relapse	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 long term	1	61	Risk Ratio (M-H, Fixed, 95% CI)	0.32 [0.05, 2.14]
2 Leaving the study early	2	145	Risk Ratio (M-H, Fixed, 95% CI)	2.38 [1.15, 4.93]
3 Death	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
3.1 long term	1	84	Risk Ratio (M-H, Fixed, 95% CI)	2.61 [0.11, 62.26]

Analysis 10.1. Comparison 10 SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus CLIENT-FOCUSED CASE MANAGEMENT, Outcome 1 Global state: Relapse.

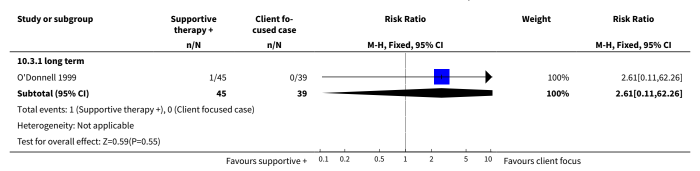




Analysis 10.2. Comparison 10 SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus CLIENT-FOCUSED CASE MANAGEMENT, Outcome 2 Leaving the study early.



Analysis 10.3. Comparison 10 SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus CLIENT-FOCUSED CASE MANAGEMENT, Outcome 3 Death.



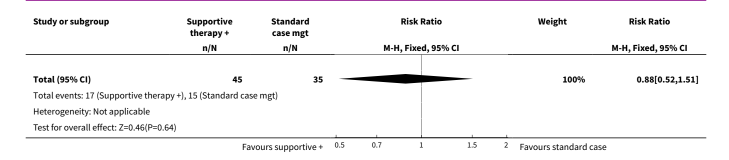
Comparison 11. SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus STANDARD CASE MANGEMENT

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Leaving the study early	1	80	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.52, 1.51]
2 Death	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 long term	1	80	Risk Ratio (M-H, Fixed, 95% CI)	2.35 [0.10, 55.94]

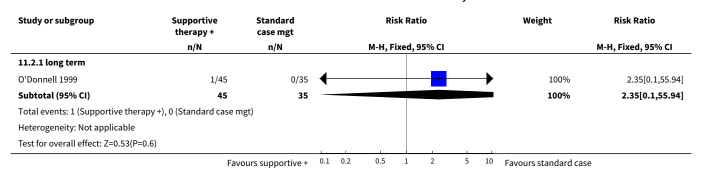
Analysis 11.1. Comparison 11 SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus STANDARD CASE MANGEMENT, Outcome 1 Leaving the study early.

Study or subgroup	Supportive therapy +	Standard case mgt	Risk Ratio				Weight	Risk Ratio	
	n/N	n/N		М-Н	, Fixed, 95%	% CI			M-H, Fixed, 95% CI
O'Donnell 1999	17/45	15/35	_		-			100%	0.88[0.52,1.51]
	Favo	urs supportive +	0.5	0.7	1	1.5	2	Favours standard case	!





Analysis 11.2. Comparison 11 SUPPORTIVE THERAPY PLUS CLIENT-FOCUSED CASE MANAGEMENT versus STANDARD CASE MANGEMENT, Outcome 2 Death.



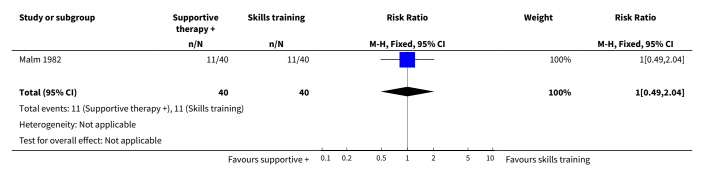
Comparison 12. SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Global state: 1. Relapse	1	80	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.49, 2.04]
2 Global state: 2. No remission	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 long term	1	80	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.54, 1.12]
3 Service outcomes: Not discharged from hospital	1	80	Risk Ratio (M-H, Fixed, 95% CI)	1.14 [0.46, 2.85]
4 Mental state: 1. Average endpoint score for 'inability to feel' (CPRS, high = poor)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
4.1 long term	1	80	Mean Difference (IV, Fixed, 95% CI)	0.10 [-0.08, 0.28]
5 Mental state: 2. Average endpoint score for 'derealisation' (CPRS, skewed data, high = poor)			Other data	No numeric data

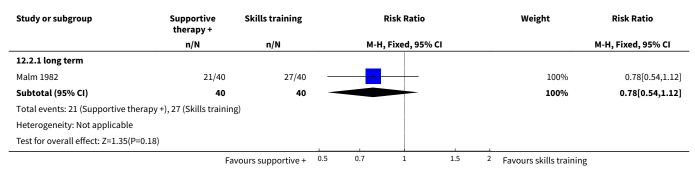


Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
5.1 long term			Other data	No numeric data
6 Leaving the study early	1	80	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.35, 2.84]
7 General functioning: 1. Average endpoint score for 'free time activities' (KAS, high = good)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
7.1 long term	1	80	Mean Difference (IV, Fixed, 95% CI)	0.10 [0.02, 0.18]
8 General functioning: 2. Average end- point score for 'withdrawal' (KAS, skewed data, high = good)			Other data	No numeric data
8.1 long term			Other data	No numeric data
9 Death	1	80	Risk Ratio (M-H, Fixed, 95% CI)	2.0 [0.19, 21.18]

Analysis 12.1. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 1 Global state: 1. Relapse.

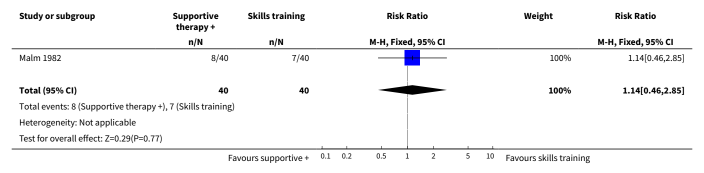


Analysis 12.2. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 2 Global state: 2. No remission.





Analysis 12.3. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 3 Service outcomes: Not discharged from hospital.



Analysis 12.4. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 4 Mental state: 1. Average endpoint score for 'inability to feel' (CPRS, high = poor).

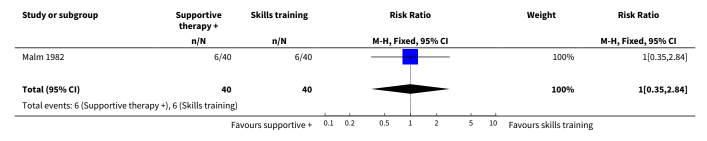
Study or subgroup	Supportive therapy +		Skills training		Mean Difference			Weight	Mean Difference	
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% CI			Fixed, 95% CI
12.4.1 long term										
Malm 1982	40	1.7 (0.4)	40	1.6 (0.4)					100%	0.1[-0.08,0.28]
Subtotal ***	40		40						100%	0.1[-0.08,0.28]
Heterogeneity: Not applicable										
Test for overall effect: Z=1.12(P=0.26)										
			Favours	supportive +	-0.5	-0.25	0 0.2	5 0.5	Favours skil	lls training

Analysis 12.5. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 5 Mental state: 2. Average endpoint score for 'derealisation' (CPRS, skewed data, high = poor).

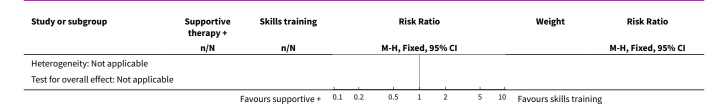
Mental state: 2. Average endpoint score for 'derealisation' (CPRS, skewed data, high = poor)

Study	Intervention	N	Me	an SD	
		long t	erm		
Malm 1982	Supportive therapy + skills training	40	0.5	0.27	
Malm 1982	Skills training	40	0.7	0.27	

Analysis 12.6. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 6 Leaving the study early.







Analysis 12.7. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 7 General functioning: 1. Average endpoint score for 'free time activities' (KAS, high = good).

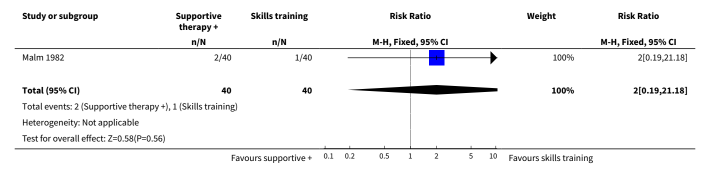
Study or subgroup		pportive erapy +	Skill	s training		Мє	ean Difference		Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)		F	ixed, 95% CI			Fixed, 95% CI
12.7.1 long term										
Malm 1982	40	0.5 (0.2)	40	0.4 (0.2)			-		100%	0.1[0.02,0.18]
Subtotal ***	40		40				•		100%	0.1[0.02,0.18]
Heterogeneity: Not applicable										
Test for overall effect: Z=2.48(P=0.01)										
			Favours	skills training	-0.5	-0.25	0 0.25	0.5	Favours sup	portive +

Analysis 12.8. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 8 General functioning: 2. Average endpoint score for 'withdrawal' (KAS, skewed data, high = good).

General functioning: 2. Average endpoint score for 'withdrawal' (KAS, skewed data, high = good)

Study	Intervention		N	Mean	SD
		lon	g term		
Malm 1982	Supportive therapy + skills training	40	0.7	0.45	
Malm 1982	Skills training	40	0.5	0.45	

Analysis 12.9. Comparison 12 SUPPORTIVE THERAPY PLUS SKILLS TRAINING versus SKILLS TRAINING, Outcome 9 Death.



ADDITIONAL TABLES



Table 1. Design of a future study

Methods	Allocation: centralised sequence generation with table of random numbers or computer-generated code, stratified by severity of illness, sequence concealed till interventions assigned. Blinding: those recruiting and assigning participants, those assessing outcomes, all blind to allocated group, blinding could be tested. Duration: minimum of 24 weeks.
Participants	Diagnosis: schizophrenia, if operational criteria used these should be in the context of routine care. N = 450*. Age: adults. Sex: men and women. Setting: anywhere.
Interventions	 Supportive therapy: the nature and frequency of this approach should be clearly described. N = 150. Routine therapy. N = 150. This also should be clearly described. Cognitive behavioural therapy. N = 150.
Outcomes	Quality of life: healthy days,** SF-36***. Service outcomes: days in hospital, time attending psychiatric outpatient clinic. Satisfaction with care: patients/carers. Global state: CGI.*** Mental state: CGI. Social functioning: to include occupational status. Adverse effects: including mortality. Economic data.
Notes	* size of study to detect a 10% difference in improvement with 80% certainty. ** Primary outcome. *** If scales are used to measure outcome then there should be binary cut off points, defined before study starts, of clinically important improvement.

N = number of participants, BPRS = Brief Psychiatric Rating Scale, CGI = Clinical Global Impressions Scale

APPENDICES

Appendix 1. Previous searches

1. Electronic searches

We searched the Cochrane Schizophrenia Group's study-based register (January 2004) using the phrase:

[((*support* OR *advoc*) in REFERENCE) and ((*support* or *individual* or *sociotherapy* or *socioenvir*) in Study)]

This register is compiled by systematic searches of major databases, hand searches and conference proceedings (see Group Module).

2. Reference searching

We inspected reference lists of all identified studies (included or excluded) for more studies.

3. Personal contact

We contacted authors of relevant reviews or studies for other sources of relevant information.

Appendix 2. Previous data analysis methods

1. Selection of trials

We (LB, TP) independently inspected the abstracts of all citations identified from the search. We identified potentially relevant abstracts and ordered full papers and reassessed these for inclusion and methodological quality. We resolved any disagreement by discussion and reported the outcome.



2. Assessment of quality

We (LB, TP) independently allocated trials to three quality categories, as described in the Cochrane Collaboration Handbook (Higgins 2005), which is based on the evidence of a strong relationship between allocation concealment and direction of effect (Schulz 1995). Where disagreement arose as to which category a trial should be allocated, we attempted to resolve this by discussion. If doubt remained we added the study to the list of trials awaiting assessment until further information could be obtained. Only trials in Category A or B were included in the review.

- A. Low risk of bias (adequate allocation concealment)
- B. Moderate risk of bias (some doubt about the results)
- C. High risk of bias (inadequate allocation concealment).

For the purpose of the analysis in this review, trials were included if they met the Cochrane Handbook criteria A or B.

- 3. Data management
- 3.1 Data extraction

We (LB, TP) independently extracted data and contacted authors' of trials to provide missing data.

3.2 Intention-to-treat analysis

We excluded data from studies where more than 50% of participants in any group were lost to follow-up (this did not include the outcome of 'leaving the study early'). In studies with less than 50% dropout rate, people leaving early were considered to have had the negative outcome, except for the event of death. The impact of including studies with high attrition rates (25% to 50%) were analysed in a sensitivity analysis. If inclusion of data from this latter group did result in a substantive change in the estimate of effect, their data were not added to trials with less attrition, but were presented separately.

- 4. Data analysis
- 4.1 Binary data

For binary outcomes we calculated the relative risk (RR) and its 95% confidence interval (CI). The number needed to treat statistic (NNT) and number needed to harm (NNH) were also calculated. If data were heterogeneous (see section 5) we used a random-effects model.

4.2 Continuous data

- 4.2.1 Skewed data: continuous data on clinical and social outcomes are often not normally distributed. To avoid the pitfall of applying parametric tests to non-parametric data the following standards were applied to all data before inclusion: (a) standard deviations and means were reported in the paper or obtainable from the authors, (b) when a scale started from a finite number (such as zero), the standard deviation, when multiplied by two, should be less than the mean (as otherwise the mean was unlikely to be an appropriate measure of the centre of the distribution (Altman 1996)). Endpoint scores on scales often have a finite start and end point and this rule can be applied to them.
- 4.2.2 Summary statistic: for continuous outcomes we estimated the weighted mean difference (WMD) between groups. Again, if data were heterogeneous (see section 5) we used a random effects model. In circumstances where trials assessed the same outcome but measured it in a variety of ways, then we analysed data using the standardised mean difference (SMD).
- 4.2.3 Valid scales: Unpublished scales are known to be subject to bias in trials of treatments for schizophrenia (Marshall 2000) therefore, we only included continuous data from rating scales were if the measuring instrument had been described in a peer-reviewed journal and the instrument was either a self report or completed by an independent rater or relative (not the therapist).
- 4.2.4 Endpoint versus change data: where possible we presented endpoint data and if both endpoint and change data were available for the same outcomes then we only reported the former in this review.
- 4.2.5 Cluster trials: studies increasingly employ 'cluster randomisation' (such as randomisation by clinician or practice) but analysis and pooling of clustered data poses problems. Authors often fail to account for intra class correlation in clustered studies, leading to a 'unit of analysis' error (Divine 1992) whereby p values are spuriously low, confidence intervals unduly narrow and statistical significance overestimated causing type I errors (Bland 1997, Gulliford 1999).

Where clustering was not accounted for in primary studies, we presented the data in a table, with a (*) symbol to indicate the presence of a probable unit of analysis error. In subsequent versions of this review we will seek to contact first authors of studies to obtain intra class correlation co-efficients of their clustered data, and to adjust for this using accepted methods (Gulliford 1999). Where clustering has been incorporated into the analysis of primary studies, we will also present these data as if from a non-cluster randomised study, but adjusted for the clustering effect.

We have sought statistical advice and have been advised that the binary data as presented in a report should be divided by a 'design effect'. This is calculated using the mean number of participants per cluster (m) and the intraclass correlation co-efficient (ICC) [Design effect = 1+(m-1)*ICC] (Donner 2002). If the ICC was not reported, it was assumed to be 0.1 (Ukoumunne 1999).



Where cluster studies were appropriately analysed taking into account intra-class correlation coefficients and relevant data documented in the report, synthesis with other studies was possible using the generic inverse variance technique.

5. Investigation for heterogeneity

Firstly, we considered all included studies within any comparison to judge clinical heterogeneity. We also inspected the graphs to investigate the possibility of statistical heterogeneity. We supplemented this using the I-squared statistic. This provides an estimate of the percentage of variability due to heterogeneity rather than chance alone. Where the I-squared estimate is equal to or greater than 75%, we interpreted this as indicating the presence of high levels of heterogeneity (Higgins 2003). We did not summate data if inconsistency were high, but presented the data separately and investigated reasons for heterogeneity.

6. Addressing publication bias

We had planned to enter data from all included studies into a funnel graph (trial effect against trial size) in an attempt to investigate the likelihood of overt publication bias (Davey 1997). However, we decided not to undertake funnel graphs, due to insufficient data being available to enable meaningful conclusions to be drawn.

7. Sensitivity analyses

We analysed the effect of including studies with high attrition rates in a sensitivity analysis.

8. General

Where possible, we entered data in such a way that the area to the left of the line of no effect indicated a favourable outcome for supportive therapy and supportive care.

WHAT'S NEW

Date	Event	Description
8 November 2017	Amended	New author team updating this review. It will be split into three new titles, creating a 'family' of Supportive Therapy reviews. New protocols to be published.

HISTORY

Protocol first published: Issue 1, 2004 Review first published: Issue 1, 2007

Date	Event	Description			
5 February 2015	New citation required but conclusions have not changed	Results from 2012 search added to review, four new studies added. No new conclusions.			
28 November 2012	New search has been performed	2012 update search run 58 new references found.			
25 April 2008	Amended	Converted to new review format.			
27 April 2007	New citation required and conclusions have changed	Substantive amendment			

CONTRIBUTIONS OF AUTHORS

Lucy Buckley - primary author for original version of review. Protocol development, main author of text of review, quality rating and selection of studies, data extraction, data entry and analysis using RevMan.

Nicola Maayan - screening, data extraction and analysis of new studies from the 2012 search, updated the results, risk of bias of all studies and 'Summary of findings' tables.



Karla Soares-Weiser - screening, data extraction and analysis of new studies from the 2012 search, contributed to 'Summary of findings' tables.

Clive Adams - protocol development, contributed to text of review, provided advice and motivation.

DECLARATIONS OF INTEREST

Lucy Buckley - none known.

Karla Soares-Weiser and Nicola Maayan - work for Enhance Review Ltd, a company that carries out systematic reviews mostly for the public sector. We currently do not provide services for the pharmaceutical industry.

Clive Adams - none known.

SOURCES OF SUPPORT

Internal sources

• Affinity Healthcare, Cheadle Royal Hospital, UK.

External sources

- Leeds Community and Mental Health Services, NHS Teaching Trust, UK.
- · Northumberland Tyne and Wear NHS Trust, UK.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The new methods template includes new methodology now required for Cochrane reviews, as such the methods section includes updates and differences from the original protocol.

The outcomes related to medication (insight, compliance, change in medication and attitude to medication) have been added to the included outcomes.

INDEX TERMS

Medical Subject Headings (MeSH)

Antipsychotic Agents [therapeutic use]; Family Therapy; Intention to Treat Analysis; Mental Health Services; Patient Satisfaction; Psychotherapy [methods]; Randomized Controlled Trials as Topic; Schizophrenia [*therapy]; Schizophrenic Psychology; Social Support

MeSH check words

Humans