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Fluphenazine (oral) versus atypical antipsychotics for schizophrenia (Review)

Sampford JR, Sampson S, Li BG, Zhao S, Xia J, Furtado VA

Sampford JR, Sampson S, Li BG, Zhao S, Xia J, Furtado VA.
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[Intervention Review]

Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

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ABSTRACT

Background

Fluphenazine is a typical antipsychotic drug from the phenothiazine group of antipsychotics. It has been commonly used in the treatment of schizophrenia, however, with the advent of atypical antipsychotic medications, use has declined over the years.

Objectives

To measure the outcomes (both beneficial and harmful) of the clinical effectiveness, safety and cost-effectiveness of oral fluphenazine versus atypical antipsychotics for schizophrenia.

Search methods

We searched the Cochrane Central Register of Studies (25 April 2013). For the economic search, we searched the Cochrane Schizophrenia Group Health Economic Database (CSzGHED) on 31 January 2014

Selection criteria

All randomised controlled trials (RCTs) comparing fluphenazine (oral) with any other oral atypical antipsychotics.

Data collection and analysis

Review authors worked independently to inspect citations and assess the quality of the studies and to extract data. For homogeneous dichotomous data we calculated the risk ratio (RR) and 95% confidence interval (CI), and calculated the mean differences (MDs) for continuous data. We assessed risk of bias for included studies and used GRADE (Grading of Recommendations Assessment, Development and Evaluation) to rate the quality of the evidence.

Main results

Four studies randomising a total of 202 people with schizophrenia are included. Oral fluphenazine was compared with oral amisulpride, risperidone, quetiapine and olanzapine.

Comparing oral fluphenazine with amisulpride, there was no difference between groups for mental state using the Brief Psychiatric Rating Scale (BPRS) (1 RCT, n = 57, MD 5.10 95% CI -2.35 to 12.55, *very low-quality evidence*), nor was there any difference in

numbers leaving the study early for any reason (2 RCTs, n = 98, RR 1.19 95% CI 0.63 to 2.28, *very low-quality evidence*). More people required concomitant anticholinergic medication in the fluphenazine group compared to amisulpride (1 RCT, n = 36, RR 7.82 95% CI 1.07 to 57.26, *very low-quality evidence*). No data were reported for important outcomes including relapse, changes in life skills, quality of life or cost-effectiveness.

Comparing oral fluphenazine with risperidone, data showed no difference between groups for 'clinically important response' (1 RCT, n = 26, RR 0.67 95% CI 0.13 to 3.35, *very low-quality evidence*) nor leaving the study early due to inefficacy (1 RCT, n = 25, RR 1.08 95% CI 0.08 to 15.46, *very low-quality evidence*). No data were reported for relapse; change in life skills; quality of life; extrapyramidal adverse effects; or cost-effectiveness.

Once again there was no difference when oral fluphenazine was compared with quetiapine for clinically important response (1 RCT, n = 25, RR 0.62 95% CI 0.12 to 3.07, *very low-quality evidence*), nor leaving the study early for any reason (1 RCT, n = 25, RR 0.46 95% CI 0.05 to 4.46, *very low-quality evidence*). No data were reported for relapse; clinically important change in life skills; quality of life; extrapyramidal adverse effects; or cost-effectiveness.

Compared to olanzapine, fluphenazine showed no superiority for clinically important response (1 RCT, n = 60, RR 1.33 95% CI 0.86 to 2.07, *very low-quality evidence*), in incidence of akathisia (1 RCT, n = 60, RR 3.00 95% CI 0.90 to 10.01, *very low-quality evidence*) or in people leaving the study early (1 RCT, n = 60, RR 3.00 95% CI 0.33 to 27.23, *very low-quality evidence*). No data were reported for relapse; change in life skills; quality of life; or cost-effectiveness.

Authors' conclusions

Measures of clinical response and mental state do not highlight differences between fluphenazine and amisulpride, risperidone, quetiapine or olanzapine. Largely measures of adverse effects are also unconvincing for substantive differences between fluphenazine and the newer drugs. All included trials carry a substantial risk of bias regarding reporting of adverse effects and this bias would have favoured the newer drugs. The four small short included studies do not provide much clear information about the relative merits or *disadvantages* of oral fluphenazine compared with newer atypical antipsychotics.

PLAIN LANGUAGE SUMMARY

Comparing effectiveness of an older antipsychotic (oral fluphenazine) with newer antipsychotics for treating schizophrenia

Introduction

People with schizophrenia often hear voices or see things (hallucinations) and have strange beliefs (delusions). It is a distressing and debilitating illness. The main treatment for schizophrenia is antipsychotic drugs. Fluphenazine is an older antipsychotic drug first formulated in the 1950s, effective for treating the psychoses of schizophrenia. However fluphenazine can cause some serious side effects, particularly movement disorders, and is known to lower people's mood. Fluphenazine is inexpensive but the arrival of newer antipsychotic drugs with fewer movement disorder side effects reduced its use and market share.

Methods

An electronic search of Cochrane Schizophrenia's register of studies was carried out in 2013. Review authors looked for trials that randomised people with schizophrenia to receive either oral fluphenazine or an atypical antipsychotic. Four studies with a total of 202 people with schizophrenia could be included. The trials compared fluphenazine with either amisulpride, risperidone, quetiapine or olanzapine.

Results

Data showed oral fluphenazine is no better or worse in improving mental state than amisulpride but more people receiving oral fluphenazine did need to take additional anticholinergic medication (drugs used to help relieve a range of symptoms such as involuntary movements of the muscles, high blood pressure and insomnia).

Data from the trials comparing oral fluphenazine with either risperidone, quetiapine or olanzapine also showed no superiority between the treatment groups for clinical improvement. Only the trial comparing oral fluphenazine with olanzapine provided adverse-effects data. Again, incidence of akathisia, a movement disorder, was similar between treatment groups.

Quality of evidence

Evidence from these few trials is poor, of low quality and involves a small number of participants. It does not provide clear overall information about whether oral fluphenazine is better or worse than atypical antipsychotic drugs for treating people with schizophrenia. Data were not available for important outcomes such as such, relapse, hospital admission, satisfaction, costs and quality of life. Adverse-effects data were poorly reported. Future large-scale research should report on these important outcomes.

Conclusions

Fluphenazine is low cost and widely available, so is likely to remain one of the most widely used treatments for schizophrenia worldwide. However, evidence currently available from randomised controlled trials about its effectiveness compared to atypical antipsychotics is unclear.

SUMMARY OF FINDINGS FOR THE MAIN COMPARISON *[Explanation]*

FLUPHENAZINE (ORAL) compared to AMISULPRIDE for schizophrenia						
Patient or population: patients with schizophrenia Settings: Austria & EU Intervention: FLUPHENAZINE (ORAL) Comparison: AMISULPRIDE						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	AMISULPRIDE	FLUPHENAZINE (ORAL)				
Mental state: Average endpoint score BPRS total score - short term (up to 12 weeks) (high = poor) Brief Psychiatric Rating Scale (BPRS). Scale from: 0 to 108. Follow-up: 3 weeks	The mean mental state: average endpoint score BPRS total score - short term (up to 12 weeks) (high = poor) in the control groups was 37.2 points	The mean mental state: average endpoint score BPRS total score - short term (up to 12 weeks) (high = poor) in the intervention groups was 5.1 higher (-2.35 to 12.55 higher)		57 (1 study)	⊕○○○ very low ^{1,2}	
Relapse (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Clinically important change in life skills (long term) - not measured	See comment	See comment	Not estimable	-	See comment	No study measured this outcome
Quality of life (long term) - not measured	See comment	See comment	Not estimable	-	See comment	No study measured this outcome

Adverse effects: Extrapyramidal effects - concomitant anticholinergic medication - short term (up to 12 weeks) Participants requiring concomitant anticholinergic medication Follow-up: 3 weeks	53 per 1000³	412 per 1000 (56 to 1000)	RR 7.82 (1.07 to 57.26)	36 (1 study)	⊕○○○ very low^{2,4}	
Leaving the study early - any reason - short term (up to 12 weeks) Follow-up: 3 weeks	Moderate		RR 1.19 (0.63 to 2.28)	98 (2 studies)	⊕○○○ very low^{2,4}	
	10 per 1000³	33 per 1000 (1 to 768)				
Cost-effectiveness (long term) - not measured	See comment	See comment	Not estimable	-	See comment	No study measured this outcome

*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: rated 'serious' - randomisation methods not clearly stated; not all outcomes reported, not all participants accounted for. Only one small study included ([Boyer 1987](#), n = 62).

² Imprecision: rated 'very serious' - few participants, few events, leading to uncertainty in the precision of estimate of effect.

³ Control risk: mean baseline risk presented from single study.

⁴ Risk of bias: rated 'serious' - randomisation methods not clearly stated; not all outcomes reported, sponsored by pharmaceutical company. Only one small study included ([Saletu 1994](#), n = 40).

BACKGROUND

Description of the condition

Schizophrenia is a psychotic disorder that can present with a variety of psychotic, cognitive and affective symptoms. It generally follows a chronic course with acute relapses and (often partial) remission. Schizophrenia is diagnosed in approximately 15.2 people per 100,000 per year (McGrath 2008). The prevalence is higher, at 4.6 per 1000 (Saha 2005), which is another sign of the chronicity of the condition. Heritability studies indicate a significant genetic component to the aetiology, however attempts to discover genes that directly cause schizophrenia have not been fruitful. Many environmental risk factors (such as urbanicity, deprivation, migrant status, fetal anoxia, childhood abuse, cannabis misuse etc.) have been shown to increase the risk of developing schizophrenia. Hence, it is currently hypothesised that the aetiology is a polygenic susceptibility to schizophrenia in individuals, which interacts with environmental risk factors. Research is increasingly focusing on these genetic and environmental interactions (van Os 2008). Symptoms are often sub-divided into 'positive' and 'negative' symptoms: positive symptoms include delusions (fixed false beliefs) and hallucinations (perceptions in the absence of an external stimulus). Negative symptoms are harder to define but often involve reductions in emotional and executive functioning, for example flattened affect, self-neglect, social isolation and apathy. Morbidity is considerable, with the majority of sufferers unable to work (Marvaha 2004). There is also increased mortality - particularly due to suicide (Healy 2012).

Description of the intervention

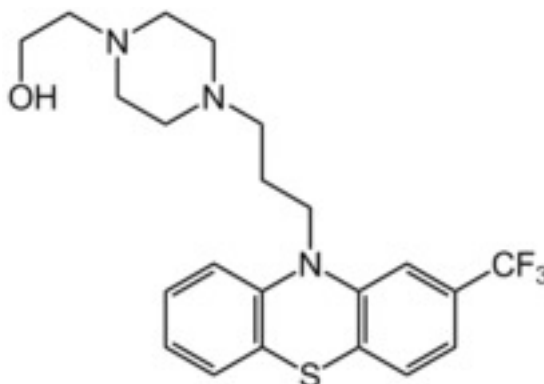
Antipsychotics are the most effective available treatment for schizophrenia and are most effective at treating the positive symptoms of schizophrenia, however they are poorer at treating the

negative symptoms (Kane 1986). Antipsychotics can be classified in a number of ways; commonly they are divided into typical and atypical groups. Fluphenazine, developed by Bristol Myers-Squibb and approved by the US Food and Drug Administration (FDA) in 1959, is a typical antipsychotic piperazine drug from the phenothiazine group of antipsychotics. It is available as a tablet, short-acting injection or long-acting injection. Originally, fluphenazine was used in Britain for the treatment of anxiety, until American reports highlighted its potential for the treatment of psychotic illness (Darling 1959; Millar 1963). Since then, it has been commonly used in the treatment of schizophrenia; it is acknowledged as an essential medicine by the World Health Organization (WHO) and widely used internationally (WHO 2005). However, with the advent of atypical antipsychotic medications, use has declined over the years.

How the intervention might work

Multiple lines of evidence point to an excess of dopaminergic neuro-transmission in schizophrenia. All antipsychotics are thought to be effective by reducing dopamine receptor activity, usually by dopamine blockade in the mesolimbic area of the brain (Grace 1991). Fluphenazine (2-[4-[3-[2-(trifluoromethyl)-10H-phenothiazin-10-yl]propyl]piperazin-1-yl]ethanol, Figure 1) is a high-potency D2 antagonist and also blocks D1a receptors postsynaptically (Seeman 2002). It is not wholly specific: this and other receptor activities account for its side-effect profile. These side effects range from hypotension secondary to alpha-adrenergic blockade, anticholinergic symptoms and extrapyramidal side effects (EPSEs) (tardive dyskinesia, muscle rigidity, tremor, dystonias and akathisia). It can also induce the neuroleptic malignant syndrome. It has variable inter-individual bioavailability and undergoes extensive first-pass metabolism. Peak plasma levels occur within hours and half-life is approximately 15 hours (Dencker 1988; Dysken 1981).

Figure 1. Fluphenazine structure



Why it is important to do this review

Recent guidelines support the use of atypical antipsychotics as the first-line treatment in schizophrenia (APA 2004). Pharmaceutical companies have marketed atypical medications as being superior to typical in terms of their efficacy and tolerability (Kendall 2011), whereas recent trials dispute this supposed advantage (Jones 2006; Leucht 2009; Lieberman 2005). It is acknowledged that typical drugs may have a higher propensity for EPSEs than many atypical drugs, many of which are more likely to induce the metabolic syndrome. However, EPSEs can often be avoided by low-dose prescribing. There are increasing concerns about the cardiovascular risks associated with long-term use of atypical antipsychotics. Additionally, it is the inexpensive typical antipsychotics that are more heavily used instead of the more expensive atypical options in the developing world. It is the accumulation of such factors that have renewed interest in researching the efficacy and tolerability of typical antipsychotics. Currently there is a lack of research evidence on fluphenazine versus atypical antipsychotics and this review aims to draw together the existing evidence.

In terms of the costs of schizophrenia, this was estimated at about £6.7 billion in England in 2004/05, of which the direct costs were £2 million while the indirect costs accounted for the rest (Mangalore 2007). The cost of fluphenazine (oral) itself is inexpensive compared to other atypical antipsychotics, at £1.88 for a 10 milligram (mg) tablet. The maximum daily dose of fluphenazine (oral) is 10 mg per day, which costs £1.88 per day, or £56.40 per month (fluphenazine oral is not present in the BNF - the cost was in US Dollars and was converted to GBP on 31st January 2014 at the prevailing exchange rate on that day). The atypical antipsychotics in comparison are more expensive than typical antipsychotics, with olanzapine available at £13.11 for 28 5 mg tablets, and clozapine (Clozaril) at £21.56 for 28 100 mg tablets.

It is important to complement the clinical effectiveness of fluphenazine (oral) with its cost-effectiveness; Davies and colleagues (Davies 2007) conducted a study on cost-effectiveness of the first-generation antipsychotics (i.e. flupentixol, trifluoperazine, chlorpromazine) and the second generation antipsychotics (i.e. risperidone, olanzapine, amisulpride). The study findings argue that there is no evidence to suggest that atypical (second generation) antipsychotics are more cost-effective than typical (first-generation) antipsychotics.

This is one of a family of related Cochrane reviews (Table 1).

OBJECTIVES

Fluphenazine (oral) versus atypical antipsychotics for schizophrenia (Review)
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To measure the outcomes (both beneficial and harmful) of the clinical effectiveness, safety and cost-effectiveness of oral fluphenazine versus atypical antipsychotics for schizophrenia.

METHODS

Criteria for considering studies for this review

Types of studies

All relevant randomised controlled trials (RCTs). We planned to include data from cross-over trials only until the point of the first cross-over as thereafter data tend to become unstable. If trials were described as 'double-blind' but implied randomisation, we included them in a sensitivity analysis (see Sensitivity analysis). We excluded quasi-randomised studies, such as those allocating by alternate days of the week. Where people were given additional treatments with oral fluphenazine, we only included data if the adjunct treatment was evenly distributed between groups and it was only the oral fluphenazine that was randomised.

With regards to selecting studies for economic evaluations, review authors (SS and VF) categorised studies as follows.

Type A - Full economic evaluation (within the framework of RCTs): studies that focus on cost-effectiveness analysis, cost-utility analysis and cost-benefit analysis.

Type B - Partial economic evaluation (within the framework of RCTs): studies that focus on cost-analysis and cost-minimisation studies of fluphenazine (oral).

Type C - Randomised trials that reported limited information, such as estimates of resources use or costs associated with fluphenazine (oral).

Types of participants

Adults (aged 18 and over) with schizophrenia or related disorders, including schizophreniform disorder, schizoaffective disorder and delusional disorder, again by any means of diagnosis. We excluded children and people with dementing illnesses, depression and primary problems associated with substance misuse. We are interested in making sure that information is as relevant to the current care of people with schizophrenia as possible so aimed to highlight clearly the current clinical state (acute, early post-acute, partial remission, remission) as well as the stage (prodromal, first episode, early illness, persistent) and whether the studies primarily focused on people with particular problems (for example, negative symptoms, treatment-resistant illnesses).

Types of interventions

1. Oral fluphenazine

Any dose or form of oral application (i.e. not depot or short-acting parenteral).

2. Atypical oral antipsychotics

Any dose or form of oral atypical antipsychotics.

Types of outcome measures

We divided outcomes into short term (up to 12 weeks), medium term (13 to 26 weeks) and long term (over 26 weeks).

Primary outcomes

1. Clinically important response (as defined by the individual studies)

1.1 Global impression - $\geq 50\%$ improvement on any relevant rating scale

Secondary outcomes

1. Death

- 1.1 Suicide
- 1.2 Natural causes

2. Global state

- 2.1 Clinically important change in global state (as defined by individual studies)
- 2.2 Average endpoint/change in global state score
- 2.3 Relapse (as defined in each study)

3. Service outcomes

- 3.1 Hospitalisation/re-hospitalisation
- 3.2 Time to hospitalisation

4. Mental state

- 4.1 Clinically important change in general mental state
- 4.2 Average endpoint/change in general mental state score
- 4.3 Clinically important change in specific symptoms (positive/negative symptoms and depression scores)

5. General functioning

- 5.1 Clinically important change in general functioning
- 5.2 Average endpoint/change in general functioning score
- 5.3 Clinically important change in specific aspects of functioning (including social skills, life skills, employment)
- 5.4 Average endpoint/change in specific aspects of functioning (including social skills, life skills, employment)

6. Quality of life

- 6.1 Clinically important change in quality of life
- 6.2 Average endpoint/change in quality of life score

7. Satisfaction with treatment

- 7.1 Clinically important change in levels of satisfaction
- 7.2 Average endpoint/change in satisfaction

8. Adverse effects - general and specific

- 8.1 Clinically important general/specific adverse effects
- 8.2 Average endpoint/change in general/specific adverse effect score

9. Extrapyramidal adverse effects

- 9.1 Any clinically significant extrapyramidal adverse effects
- 9.2 Any clinically significant extrapyramidal side effects (EPSEs) - as defined by each study
- 9.3 Average score/change in EPSEs
- 9.4 Incidence of use of antiparkinson drugs
- 9.5 Dystonia
- 9.6 Akathisia
- 9.7 Akinesia

10. Leaving the study early - any reason

- 10.1 Leaving the study early - due to inefficacy of the intervention
- 10.2 Leaving the study early due to side effects

11. Economic outcomes

- 11.1 Average change in total cost of medical and mental health care
- 11.2 Total indirect and direct costs
- 11.3 Direct resource use:
 - 11.3.1 *Outpatients - number of contacts (GP consultation, psychiatrist, psychologists, psychiatric nurse, counsellor, social worker)*
 - 11.3.2 *Hospitalisation (taking battery of tests, patients' physical, psychiatric and psychological profile and psychological assessment, number of days, relapse)*
 - 11.3.3 *Medication (different types of antipsychotics to include dose and frequency, treatment of side effects)*

11.3.4 Psychological therapies (different types of psychological therapies to include session numbers and frequency)

11.3.5 Other resources (day centres, night shelter) and transportation for medical care visits

11.4 Indirect resource use:

11.4.1 Family, relative and friends resources

11.4.2 Police, criminal justice system

11.4.3 Benefits paid, social security payments

11.4.4 Employment agency workers, absence from work, loss of productivity

11.5 Cost-effectiveness ratios represented by the incremental cost-effectiveness ratio (ICER)

11.6 Cost-utilities represented by incremental costs per quality-adjusted life year (QALY) or disability adjusted life year (DALY)

11.7 Cost-benefit represented by net Benefit Ratio, others

12. 'Summary of findings' table

We used the GRADE approach to interpret findings (Schünemann 2008) and used GRADE profiler (GRADEPRO) to import data from RevMan 5 (Review Manager) 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' table:

1. Clinically important response in mental state (short, medium and long term)
2. Relapse (long term)
3. Clinically important change in life skills (long term)
4. Quality of life (long term)
5. Adverse effects, e.g. EPSEs (medium term)
6. Leaving the study early: any reason (medium term)
7. Cost-effectiveness (long term)

Search methods for identification of studies

Electronic searches

We searched the Central Register of Studies (25 April 2013) using the phrase:

("*clozapin*":TI OR "*clozaril*":TI OR "*leponex*":TI OR "*aripiprazole*":TI OR "*olanzapin*":TI OR "*lanzac*":TI OR "*zyprex*":TI OR "*quetiapin*":TI OR "*seroquel*":TI OR "*risperidon*":TI OR "*belivon*":TI OR "*risperdal*":TI OR "*risperin*":TI OR "*rispolin*":TI OR "*sertindol*":TI OR "*serdolect*":TI OR "*ziprasidon*":TI OR "*zotepin*":TI OR "*lodopin*":TI OR "*nipolept*":TI OR "*zopite*":TI OR "*setous*":TI OR "*majorpin*":TI OR "*remox-

iprid*":TI OR "*roxiam*":TI OR "*remidon*":TI OR "*iloperidon*":TI OR "*clozapin*":AB OR "*clozaril*":AB OR "*leponex*":AB OR "*aripiprazole*":AB OR "*olanzapin*":AB OR "*lanzac*":AB OR "*zyprex*":AB OR "*quetiapin*":AB OR "*seroquel*":AB OR "*risperidon*":AB OR "*belivon*":AB OR "*risperdal*":AB OR "*risperin*":AB OR "*rispolin*":AB OR "*sertindol*":AB OR "*serdolect*":AB OR "*serlect*":AB OR "*ziprasidon*":AB OR "*zotepin*":AB OR "*lodopin*":AB OR "*nipolept*":AB OR "*zopite*":AB OR "*setous*":AB OR "*majorpin*":AB OR "*remoxiprid*":AB OR "*roxiam*":AB OR "*remidon*":AB OR "*iloperidon*":AB OR "*clozapine*" null "*clozapin*" OR "*clozaril*" OR "*leponex*" null "*aripiprazole*" OR "*olanzapin*" OR "*lanzac*" OR "*zyprex*" OR "*quetiapin*" OR "*seroquel*" OR "*risperidon*" OR "*belivon*" OR "*risperdal*" OR "*risperin*" OR "*rispolin*" OR "*sertindol*" OR "*serdolect*" OR "*serlect*" OR "*ziprasidon*" OR "*zotepin*" OR "*lodopin*" OR "*nipolept*" OR "*zopite*" OR "*setous*" OR "*majorpin*" OR "*remoxiprid*" OR "*roxiam*" OR "*remidon*" OR "*iloperidon*" OR "*atypical*":TI OR "*atypical*":TI OR "*atypical*":AB OR "*atypical*") AND ("*fluphen*":TI OR "*fluphen*":TI OR "*flufen*":TI OR "*flufen*":TI OR "*lyogen*":TI OR "*lyogen*":TI OR "*prolixin*":TI OR "*prolixin*":TI OR "*siqualon*":TI OR "*siqualon*":TI OR "*modec*":TI OR "*moditen*":TI OR "*fluphen*":AB OR "*flufen*":AB OR "*lyogen*":AB OR "*prolixin*":AB OR "*siqualon*":AB OR "*modec*":AB OR "*moditen*":AB OR "*fluphen*" OR "*flufen*" OR "*lyogen*" OR "*prolixin*" OR "*siqualon*" OR "*modec*" OR "*moditen*")

2. Economic study search of Cochrane Schizophrenia Group Health Economic Database (2013)

For the economic search, we replicated the above strategy in the Cochrane Schizophrenia Group Health Economic Database (CSzGHED) on 31 January 2014. The database of studies relates to cost-effectiveness of schizophrenia treatments. This database was constructed from systematic searches of four databases: Health Economic Evaluation Database (HEED), National Health Services Health Economic Database (NHS EED), Cost-Effectiveness Analysis Registry (CEA) and EconLit as well as Cochrane Registry.

Searching other resources

1. Reference searching

We inspected references of all included studies for further relevant studies.

2. Personal contact

We contacted the first author of each included study for information regarding unpublished trials.

3. Pharmaceutical companies

We contacted relevant pharmaceutical companies to obtain more information or data on unpublished trials if appropriate.

Data collection and analysis

Selection of studies

Review author JS independently inspected citations from the searches and identified relevant abstracts. Review author SS independently re-inspected a random 20% sample to ensure reliability. Where disputes arose, we acquired the full report for more detailed scrutiny. JS obtained and inspected full reports of the abstracts meeting the review criteria. Again, SS re-inspected a random 20% of full reports in order to ensure reliable selection. Where it was not possible to resolve disagreement by discussion, we attempted to contact the authors of the study for clarification.

For the selection of economic studies, review authors VF and SS inspected all retrieved citations identified by the economic database search, and where disputes arose, we acquired the full report for further inspection.

Data extraction and management

1. Extraction

Review author BGL, SZ, JX, independently extracted data from included studies, and SS made a random 20% check to ensure reliability. Again, we discussed any disagreement. We extracted data presented only in graphs and figures whenever possible, but included the data only if the two review authors independently had the same result.

For the economic analysis, had Type A and B studies been identified (see [Types of studies](#)), review authors VF and SS would have investigated whether appraisal had already been undertaken by [NHS EED](#) using their search tool derived for this purpose. If appraisal had not been undertaken, VF and SS would have applied the [NHS EED](#) tool to the data. For Type C studies, we planned to extract outcome data directly from the already-included effectiveness studies.

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 were described in a peer-reviewed journal ([Marshall 2000](#)); and
- b) the measuring instrument had 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 and noted this in the [Description of studies](#) section.

2.3 Endpoint versus change data

There are advantages to 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. We combined endpoint and change data in the analysis as we preferred mean differences (MD) rather than standardised mean differences (SMD) throughout ([Higgins 2011](#)).

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 relevant data before inclusion.

Studies, $N > 200$

We entered useable data from studies of at least 200 participants, for example, in the analysis irrespective of the following rules, because skewed data pose less of a problem in large studies.

Change data

We also entered all useable change data as 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.

Endpoint data, $N < 200$

(a) When a scale started from the finite number zero, we subtracted the lowest possible value from the mean and divided this by the standard deviation (SD). If this value was lower than 1, it would have strongly suggested a skew and we excluded these data. If this ratio was higher than 1 but below 2, there is a suggestion of

skew. We entered these data and tested whether their inclusion or exclusion changed the results substantially. Finally, if the ratio was larger than 2 we included these data, because skew was less likely (Altman 1996; Higgins 2011).

b) 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.

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 PANSS (Kay 1986), this could be considered as a clinically significant response (Leucht 2005; Leucht 2005a).

2.7 Direction of graphs

We entered data in such a way that the area to the left of the line of no effect indicates a favourable outcome for oral fluphenazine. Where keeping to this makes it impossible to avoid outcome titles with clumsy double-negatives (e.g. 'Not un-improved') we reported data where the left of the line indicates an unfavourable outcome. We noted this in the relevant graphs.

Assessment of risk of bias in included studies

Again BGL and JS worked independently to assess risk of bias by using criteria described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011) to assess trial quality. This 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.

If the raters disagreed, we made the final rating by consensus with SS.

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

This review also aimed to assess the overall methodological quality of each study included in the economic evaluation. We planned to

use the checklist developed by Drummond 1996 and the CHEC criteria list (Evers 2005) for Type A and B studies. Had we found any economic studies of Type A or B level, this would have been noted in the summary as well as in a separate table.

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 (NNTB/NNTH) 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' table/s, where possible, we calculated illustrative comparative risks.

2. Continuous data

For continuous outcomes we estimated the mean difference (MD) between groups. We preferred not to calculate effect size measures (standardised mean difference (SMD)). However, if in future versions of this review, if scales of very considerable similarity are used, we will presume there is a small difference in measurement, and we will calculate effect size and transform 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. Firstly, 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 identify any cluster-randomised studies; however, in future version of this review, and where we identify studies that have not accounted for clustering in primary studies, we will present 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 coefficients (ICCs) for their clustered data and to adjust for this by using accepted methods (Gulliford 1999).

Where clustering has been incorporated into the analysis of primary studies, we will present these data as if from a non-cluster randomised study, but adjust 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 would 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). We did not identify any cross-over studies; however, in future versions of this review where such studies are identified, as both effects are very likely in severe mental illness, we will only use data from 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 and combined these 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 Systematic Reviews of Interventions* (Higgins 2011).

Dealing with missing data

1. Overall loss of credibility

At some degree of loss to 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 planned to address this within the 'Summary of findings' table/s by downgrading quality. Finally, we also planned to downgrade quality within the 'Summary of findings' table/s should loss be 25% to 50% in total. Such high losses were not experienced in the included studies.

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 are 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 stay 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 data only from people who completed the study to that point 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 was between 0% and 50%, and data only from people who completed the study to that point were reported, we reproduced these data.

3.2 Standard deviations

If standard deviations (SDs) were not reported, first, we 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 are available for group means, and either P value or 't' value are available for differences in mean, we can calculate them according to the rules described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). When only the SE is reported, SDs are calculated by the formula $SD = SE * \text{square root}(n)$. Chapters 7.7.3 and 16.1.3 of the *Cochrane Handbook for Systematic Reviews of Interventions* (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 did not impute any values, since we did not identify any missing SDs in the included studies.

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. Had such situations or participant groups arisen, we would have fully discussed these. However, meta-analysis was not possible, since all included studies compared various different atypical antipsychotics versus oral fluphenazine.

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 I^2 statistic

We investigated heterogeneity between studies by considering the I^2 statistic alongside the Chi^2 test P value. The I^2 statistic provides an estimate of the percentage of inconsistency thought to be due to chance (Higgins 2003). The importance of the observed value of I^2 depends on i) the magnitude and direction of effects and ii) the strength of evidence for heterogeneity (e.g. P value from Chi^2 test or a confidence interval for I^2). We planned to interpret an I^2 estimate greater than or equal to around 50%, accompanied by a statistically significant Chi^2 test, as evidence of substantial levels of heterogeneity (Section 9.5.2 - Higgins 2011). Had substantial levels of heterogeneity been found for the primary outcome, we would have explored the reasons for this (Subgroup analysis and investigation of heterogeneity).

Assessment of reporting biases

1. Protocol versus full study

Reporting biases arise when the dissemination of research findings is influenced by the nature and direction of results. These are described in section 10.1 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We made attempts to locate protocols for the included randomised trials. Had any protocols been available, we would have compared the outcomes in the protocol and in the published report.

2. Funnel plot

Reporting biases arise when the dissemination of research findings is influenced by the nature and direction of results (Egger 1997). These are again described in section 10 of the *Cochrane Handbook for Systematic Reviews of Interventions* (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 since there were less than 10 included studies.

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. For this review, we chose a random-effects model for all analyses.

Handling of economic data

“It has been argued for many years that promoting effective care without taking into account the cost of care and the value of any health gain can lead to inefficient use of public and private funds allocated to health care, which may indirectly result in harm for individuals and the public” (Williams 1987).

We intended to summarise data from type A and type B studies and summarise data according to the Cochrane Campbell Economic Methods Group (Higgins 2011), and if information had been available, a narrative abstract would have been presented for each included study.

We anticipated that most studies would be Type C level of economic evidence and that we would use data from such studies to calculate a GBP value associated with the outcomes. These approximate values can be calculated by:

(a) using the Personal Social Services Research Unit (PSSRU - NHS reference costs for mental health services) calculation of £338 (weighted mean average of all adult mental health in-patient bed days) per hospital bed day based in a UK NHS setting (PSSRU 2012); and

(b) assuming that one relapse equals one hospital admission, a median length of stay as per Hospital Episode Statistics 2012 (HES 2012; main speciality ‘adult mental illness’), we could utilise results of the effects of the intervention that present service use data for an adult ward as well as for relapse rates (HES is a data warehouse containing details of all admissions, outpatient appointments and A&E attendances at NHS hospitals in England);

(c) in terms of use of adjunctive medication, if the specific drug is not mentioned then we would assume that the adjunctive medication used was phenobarbital and that it would be prescribed for no longer than 14 days at an average dose of 120 mg per day; the cost for this was obtained from the BNF which provides unit costs for the medication;

(d) in terms of treatment for EPSEs, if the specific drug was not mentioned, we would assume that for the treatment procyclidine was used at a dose of 10 mg three times a day for 14 days; the cost for this was obtained from the BNF which provides unit costs for the medication

(e) in terms of treatment for akathisia, if the specific drug was not mentioned, we would assume that for the treatment propranolol was prescribed at a dose of 80 mg twice a day for 14 days; the cost for this was obtained from the BNF which provides unit costs for the medication;

(f) in terms of treatment for depression, if the specific drug was not mentioned, we would assume that for the treatment fluoxetine was prescribed at a dose of 20 mg once a day for 120 days; the cost for this was obtained from the BNF which provides unit costs for the medication;

(g) in terms of epileptic fits, we would assume that such fits last for less than five minutes (more than five minutes constitutes Status Epilepticus as specified by [NICE 2012](#)), unless otherwise specified;

(h) in terms of treatment for agitation, if the specific drug was not mentioned, we would assume that for the treatment lorazepam was prescribed at a dose of 1 mg up to four times a day for three days; the cost for this was obtained from the BNF which provides unit costs for the medication.

We did not factor any associated costs (including cost and resource use of treatment) prior to the relevant measured outcomes being considered. We are using UK NHS PSSRU reference costs of 2012 as well as BNF costs from 2013 and therefore planned to present the outcomes in terms of a GBP saving using relative risks obtained from the effectiveness part of the review, which we have considered to be a proxy for resource use.

The authors wish to emphasise the numerous assumptions that have been made for the purposes of presenting economic data, specifically of Type C studies:

1. the current included studies contributing to the Type C studies were undertaken between the years of 1987 to 2005; and, taking this into account;

2. the median length of stay and costs have been calculated from current available data, that is, according to 2012 HES costs, from primarily a UK NHS perspective; and

3. the GBP value data that are presented reflect a proxy measure only; that is, the GBP value of the intervention effect on the measured outcome, and not taking into account any costs or resource use that may likely have been incurred prior to the actual outcome (which includes, but is not limited to, costs and resource use prior to intervention, the intervention itself and post-intervention up to outcome).

We are aware that Cochrane systematic reviews are international in context and in their understanding; however, we have adopted a UK NHS perspective for the purposes of this review - partly because we have been funded by the National Institute of Health Research (NIHR) (NIHR Cochrane Programme Grant 2011, UK Reference number: 10/4001/15) to undertake a series of economic evaluations within systematic reviews.

"...[I]n the face of scarce resources, decision makers often need to consider not only whether an intervention works, but also whether its adoption will lead to a more efficient use of resources" ([Higgins 2011](#)).

Subgroup analysis and investigation of heterogeneity

1. Subgroup analyses

1.1 Primary outcomes

We subgrouped analyses by length of treatment (short, medium and long term).

1.2 Clinical state, stage or problem

Where possible, we reported data on subgroups of people in the same clinical state, stage and with similar problems.

2. Investigation of heterogeneity

Had inconsistency been high, we would have reported this. Should this happen in future versions of this review, first we will investigate whether the data have been entered correctly. Second, if the data are correct, we will visually inspect the graph and successively remove outlying studies to see if homogeneity is restored. For this review, we decided that should this occur with data contributing no more than around 10% of the total weighting to the summary finding, we will present the data. If not, we will not pool the data and discuss the issues. We know of no supporting research for this 10% cut-off but are investigating use of prediction intervals as an alternative to this unsatisfactory state.

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

Sensitivity analysis

1. Implication of randomisation

We included trials in a sensitivity analysis if they were described in some way as to imply randomisation. For the primary outcomes

we included these studies, If their inclusion did not result in a substantive difference, they remained in the analyses. If their inclusion did result in important, clinically significant but not necessarily statistically significant differences, we did not add the data from these lower quality studies to the results of the better trials, but presented such data within a subcategory.

2. Assumptions for lost binary data

Where assumptions had to be made regarding people lost to follow-up (see [Dealing with missing data](#)), we compared the findings of the primary outcomes when we use 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 the results and discussed them but continued to employ our assumption.

Where assumptions had to be made regarding missing SD 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. We undertook a sensitivity analysis 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 the results and discussed them but continued to employ our assumption.

3. Risk of bias

We analysed the effects of excluding trials that are judged to be at high risk of bias across one or more of the domains of randomisation, 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

Had we imputed any values, we would have carried out a sensitivity analysis to assess the effects of including data from trials where we used imputed values for ICC in calculating the design effect in

cluster-randomised trials. We will undertake this sensitivity analysis in future versions of this review where such imputations may be made.

If substantial differences are noted in the direction or precision of effect estimates in any of the sensitivity analyses listed above, we will not pool data from the excluded trials with the other trials contributing to the outcome, but will present them separately.

5. Fixed-effect and random-effects

We synthesised all data using a random-effects model, however, we also synthesised data for the primary outcome using a fixed-effect model to evaluate whether this alters the significance of the results.

6. Economic summary

We undertook a sensitivity analysis taking into account both the upper and lower confidence intervals for the risk ratios, of the outcomes of interest, and calculated a saving based on these values to investigate how far this affects the direction of the estimated value.

RESULTS

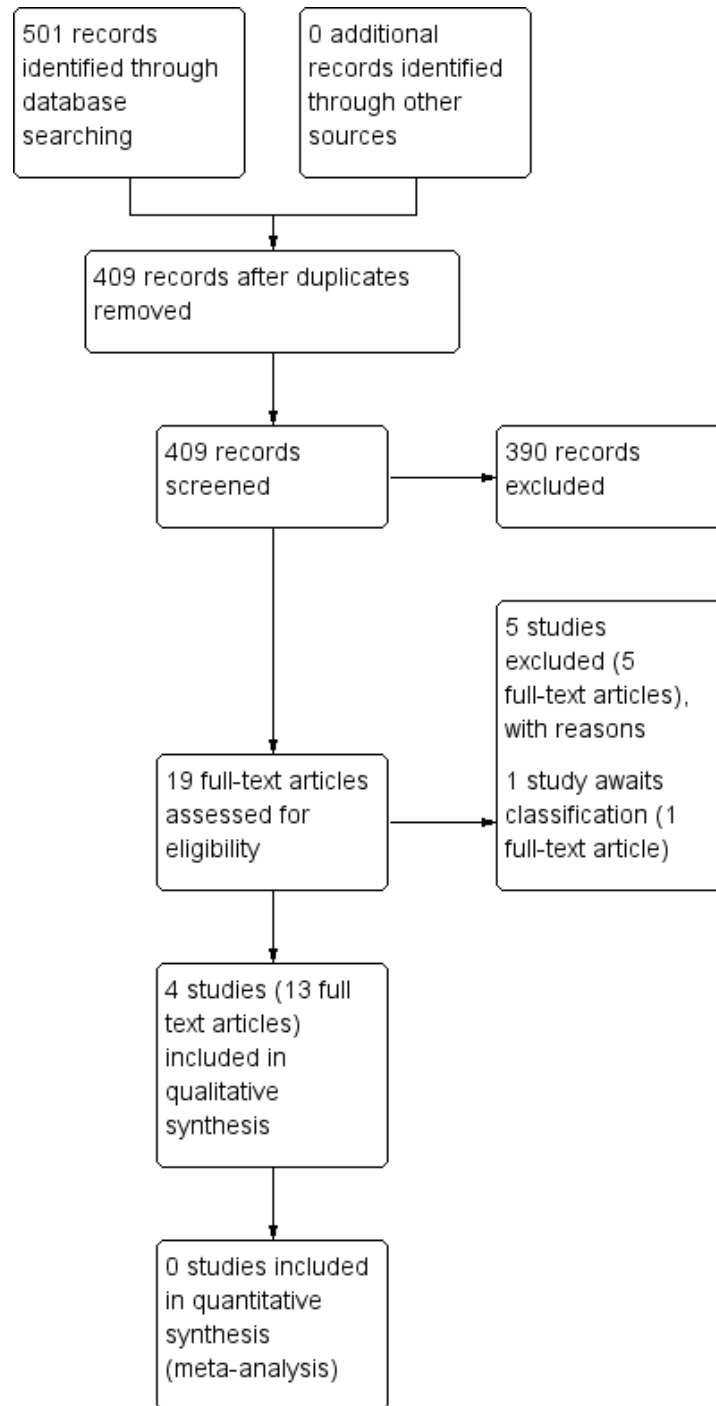
Description of studies

For substantive descriptions of studies please see [Characteristics of included studies](#), [Characteristics of excluded studies](#) and [Characteristics of studies awaiting classification](#) tables.

Results of the search

Please see [Figure 2](#) for a visual description of the study search process and study inclusion/exclusion details. Our study search identified 501 records; after duplicates were removed, we screened a total of 409 references. Of these, we excluded 390 based on title and abstract, with only 19 full-text references requested for full inspection. Of these, four studies were included.

Figure 2. Study flow diagram: 2013 study search



Included studies

1. Length of trials

Studies ranged from six weeks duration of treatment to 22 weeks. [Boyer 1987](#) and [Saletu 1994](#) both had a washout period of three weeks and three days respectively, with a treatment duration of six weeks. [Conley 2005](#) had a four- to six-week open-label lead-in phase, with a 12-week treatment duration, and [Dossenbach 1998](#) had two treatment phases, one for 'acute' (six weeks) and one for 'long term' (22 weeks).

2. Design

All included studies were parallel arm RCTs; only one study had more than two treatment arms ([Conley 2005](#)). No included study adequately described the randomisation methods used.

3. Participants

All participants had a diagnosis of schizophrenia with either DSM-III (Diagnostic and Statistical Manual, third edition) ([Boyer 1987](#); [Conley 2005](#); [Saletu 1994](#)) or DSM-IV (Diagnostic and Statistical Manual, fourth edition) ([Dossenbach 1998](#)). Participants included in [Conley 2005](#) were defined as 'treatment-refractory', and participants in [Dossenbach 1998](#) were assessed both in the 'acute' stage (with results up to six weeks), as well as the long-term treatment (up to 22 weeks).

4. Setting

Three out of the four studies provided details as to trial setting: [Conley 2005](#) was undertaken in the USA; [Dossenbach 1998](#) was undertaken in Croatia with a multicentre design; and [Saletu 1994](#) was undertaken in Austria. [Boyer 1987](#) provided no details.

5. Study size

Study sizes ranged from $n = 40$ ([Conley 2005](#); [Saletu 1994](#)) to $n = 62$ ([Boyer 1987](#)). The total number of included participants in this review is $n = 202$.

6. Interventions

6.1 Fluphenazine

The total number of participants receiving fluphenazine was $n = 92$. Doses of fluphenazine were relatively uniform between studies,

with one study permitting a larger dose range ([Dossenbach 1998](#)). [Boyer 1987](#) used a range of 2 mg to 12 mg/day; [Conley 2005](#) used a mean of 13.2 mg/day; [Dossenbach 1998](#) used a range of 5 mg to 20 mg/day, with a mean dose of 11.7 mg/day overall in both the 'acute' and 'long-term' phase of the study; [Saletu 1994](#) used a range of 2 mg to 4 mg/day.

6.2 Amisulpride

Two studies compared amisulpride with fluphenazine; the total number of participants receiving amisulpride was $n = 53$. [Boyer 1987](#) used a range of 50 mg to 300 mg/day; and [Saletu 1994](#) used a range of 50 mg to 100 mg/day.

6.3 Olanzapine

One study compared the olanzapine with fluphenazine; the total number of participants receiving olanzapine was $n = 30$. [Dossenbach 1998](#) used a range of 6 mg to 21 mg/day, with a mean average of 13.6 mg/day in the 'acute' phase, and 14.8 mg/day in the 'long-term' study phase.

6.4 Quetiapine

One study compared the quetiapine with fluphenazine; the total number of participants receiving quetiapine was $n = 12$. [Conley 2005](#) used a mean dose of 463.6 mg/day.

6.5 Risperidone

One study compared the risperidone with fluphenazine; the total number of participants receiving risperidone was $n = 13$. [Conley 2005](#) used a mean dose of 4.31 mg/day.

7. Outcomes

7.1 General remarks

We did not conduct a meta-analysis as the four included studies were presented in four different comparisons. Studies were generally lacking that compared fluphenazine oral with other atypical antipsychotics, and as a consequence, outcome-reporting between studies was not consistent. Only two studies provided data for our primary outcome of 'clinically important response' ([Conley 2005](#); [Dossenbach 1998](#)).

7.2 Acceptability and efficacy

Each included study provided data regarding mental and global state outcomes (widely-accepted rating scales, including the Brief Psychiatric Rating Scale (BPRS), Positive and Negative Symptom Scale (PANSS) and Clinical Global Impression (CGI)), however some of these data were skewed and are presented in an additional table.

7.3 Adverse events

Adverse events, including anticholinergic effects, central nervous system effects, gastrointestinal effects and 'others' were generally well-reported in the included studies. However data were seriously lacking for extrapyramidal adverse effects.

7.4 Outcome scales

7.4.1 Global state

i) Clinical Global Impression - CGI (Guy 1976)

This is a rating instrument that enables clinicians to quantify severity of illness and overall clinical improvement during therapy. A seven-point scoring system is usually used with low scores indicating decreased severity and/or greater recovery. Three studies reported data using this scale (Conley 2005; Dossenbach 1998; Saletu 1994).

7.4.2 Mental state

i) Association for Methodology and Documentation in Psychiatry - AMDP (Gebhardt 1983)

The AMDP consists of a glossary of psychopathological symptoms, as well as rating criteria to assist standardisation in recording. One included study measured degrees of apathy in participants using the AMDP manual criteria (Saletu 1994).

ii) Brief Psychiatric Rating Scale - BPRS (Overall 1962)

This scale is used to assess the severity of abnormal mental states. The original scale has 16 items, but a revised 18-item scale is commonly used. Each item is defined on a seven-point scale varying from 'not present' to 'extremely severe', scoring from zero to six or one to seven. Scores can range from zero to 108 or 18 to 126, respectively. High scores indicate more severe symptoms. The BPRS-positive cluster comprises four items, which are conceptual disorganisation, suspiciousness, hallucinatory behaviour and unusual thought content. The BPRS-negative cluster comprises only three items, which are emotional withdrawal, motor retardation, and blunted affect. Three studies reported data using this scale (Boyer 1987; Conley 2005; Dossenbach 1998).

iii) Hamilton Anxiety Scale - HAMA (Maier 1988)

HAMA is a rating scale developed to quantify the severity of anxiety symptomatology and consists of 14 items, each defined by a series of symptoms. Each item is rated on a five-point scale, ranging from zero (= not present) to four (= severe). One study reported continuous data using this scale (Dossenbach 1998).

iv) Positive and Negative Symptom Scale - PANSS (Kay 1987)

The positive and negative syndrome scale was originated as a method for evaluating positive, negative and other symptom dimensions in schizophrenia. The scale has 30 items, and each item can be rated on a seven-point scoring system varying from one (absent) to seven (extreme). This scale can be divided into three subscales for measuring the severity of general psychopathology, positive symptoms (PANSS-P) and negative symptoms (PANSS-N). A low score indicates low levels of symptoms. One study provided data using this scale (Dossenbach 1998).

v) Scale for Assessment of Negative Symptoms - SANS (Andreasen 1982)

The SANS measures the incidence and severity of negative symptoms using a 25-item scale, using a six-point scoring system, where zero = better to five = worse, where a higher score equals a more severe experience of negative symptoms. One study reported data using this scale (Saletu 1994).

7.4.3 Satisfaction with treatment

i) Drug Attitude Inventory - DAI (Hogan 1983)

The DAI is a self-administered rating scale designed to gain understanding of patient-use and personal experiences of using psychiatric medication. There are 30 items, which are rated as either 'true' or 'false' by users, including statements such as 'medication is a slow-acting poison', or 'I can't concentrate on anything when I'm on medication'. One study provided continuous data using this scale (Dossenbach 1998).

7.4.4 Adverse events

i) Assessment of Involuntary Movements Scale - AIMS (Guy 1976a)

This scale measures the examination of involuntary movements (tardive dyskinesia) consisting of 12 items scored from zero = none to four = severe, quantifying the severity of tardive dyskinesia. This scale used in short-term trials may also help to assess Parkinsonian symptoms such as tremor. One study reported continuous data using this scale (Dossenbach 1998).

ii) Hillside Akathisia Scale - HAS (Fleischhacker 1989)

The HAS was used to measure akathisia; the subjective subscale has two subjective and three objective items for which anchored rating points are provided. The subjective items take into account a patient's sensation of restlessness and urge to move, and the objective items assess physical signs of akathisia present in the head, trunk, hands, arms, feet and legs. There are a total of five items, which are measured on a five-point scoring system from zero = absent to four = present and not controllable. One study provided data using this scale (Dossenbach 1998).

iii) Leeds Sleep Evaluation Questionnaire - LSEQ (Parrott 1980)

The LSEQ is a 10-item, self-rating measurement designed to assess changes in sleep quality over the course of psychopharmacological treatment. Four domains are rated, including 'ease of initiating sleep', 'quality of sleep', 'ease of waking' and 'behaviour following wakefulness'. One study reported data using this scale (Dossenbach 1998).

iv) Simpson-Angus Scale - SAS (Simpson 1970)

The SAS measures drug-related extrapyramidal symptoms; it is a 10-item rating scale, with a score range of zero (= not present) to 40 (= severe); it includes items such as gait, rigidity, tremor and salivation. One study reported data using this scale (Dossenbach 1998).

7.5 Missing outcomes

The four included studies failed to report several of our pre-specified secondary outcomes of interest, including economic outcomes, quality of life outcomes, service-use and hospitalisation outcomes, relapse, and general function (such as social skills, employability). These are particularly patient-important outcomes that have been overlooked, and would add to the body of evidence regarding acceptability of treatment.

Excluded studies

We excluded five studies. Three, or perhaps two, studies compared amisulpride with placebo, haloperidol, or at different doses (Boyer 1986; Boyer 1987a; Boyer 1996). Pickar 1992 was not a randomised study and Ravanic 1996 provided no useable data.

Studies awaiting assessment

One study awaits assessment as only a conference abstract is available with no usable data available; the full report is required (Djukic-Dejanovic 2002).

Ongoing studies

We identified no ongoing studies.

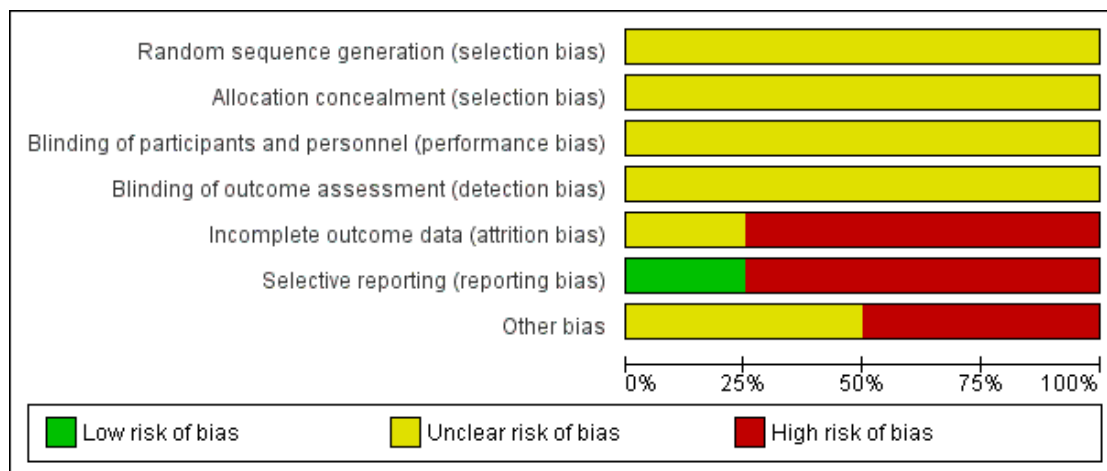
Risk of bias in included studies

For a graphical overview of 'Risk of bias' assessments in included studies, see Figure 3; Figure 4.

Figure 3. 'Risk of bias' summary: review authors' judgements about each risk of bias item for each included study.

	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
Boyer 1987	?	?	?	?	-	-	?
Conley 2005	?	?	?	?	-	-	?
Dossenbach 1998	?	?	?	?	-	+	-
Saletu 1994	?	?	?	?	?	-	-

Figure 4. 'Risk of bias' graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.



Allocation

None of the included studies provided adequate details as to randomisation methods and were all rated as an 'unclear' risk of bias. [Conley 2005](#) stated that randomisation was performed by the dispensing pharmacy; [Dossenbach 1998](#) stated that randomisation was undertaken in a 1:1 ratio; while [Boyer 1987](#) and [Saletu 1994](#) simply stated that participants were 'randomly allocated', with no further details.

Blinding

Again, none of the included studies provided adequate details as to blinding methods and were all rated as an 'unclear' risk of bias, with all studies only stating that studies were double-blinded.

Incomplete outcome data

Three included studies were rated as a 'high' risk of bias for attrition; in [Boyer 1987](#), not all participants completed ratings for various BPRS components, and it was unclear whether last observation carried forward (LOCF) was used. Forty participants were randomised in [Conley 2005](#), however data for n = 2 were 'lost', and only n = 38 (out of n = 40 randomised) were presented in the data and analysis. In [Dossenbach 1998](#), all participants were included in the safety analysis. However for efficacy n = 5 were excluded because they did not meet inclusion criteria for BPRS or CGI.

Selective reporting

Three studies were rated as a 'high' risk of bias for selective reporting; [Boyer 1987](#), [Conley 2005](#) and [Saletu 1994](#) did not report all stated outcome measures, particularly relating to continuous data with means and standard deviations not transparently reported. [Dossenbach 1998](#) was rated as a 'low' risk due to higher standards of reporting outcome data.

Other potential sources of bias

Two studies were rated as 'unclear' for other bias ([Boyer 1987](#); [Conley 2005](#)), while the other two studies rated as 'high' ([Dossenbach 1998](#); [Saletu 1994](#)). We did not detect any obvious other sources of bias with [Boyer 1987](#); study medications were supplied by Janssen Pharmaceutica and Astra-Zeneca Pharmaceuticals in [Conley 2005](#). For the two studies rated as a 'high' risk ([Dossenbach 1998](#); [Saletu 1994](#)), both were sponsored by the pharmaceutical industry, including Eli Lilly and Company ([Dossenbach 1998](#)) and Synthelabo Recherche/Laboratoires Delagrangre (Bagneux, France) ([Saletu 1994](#)).

Effects of interventions

See: [Summary of findings for the main comparison FLUPHENAZINE \(ORAL\) compared to AMISULPRIDE for schizophrenia](#); [Summary of findings 2 FLUPHENAZINE](#)

(ORAL) compared to RISPERIDONE for schizophrenia; [Summary of findings 3](#) FLUPHENAZINE (ORAL) compared to QUETIAPINE for schizophrenia; [Summary of findings 4](#) FLUPHENAZINE (ORAL) compared to OLANZAPINE for schizophrenia

COMPARISON 1: FLUPHENAZINE (ORAL) versus AMISULPRIDE

1.1 Global state: 1. Average endpoint score of CGI scales (high = poor)

1.1.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 36) ([Saletu 1994](#)). There was no significant difference between fluphenazine (oral) and amisulpride (mean difference (MD) -0.34 95% confidence interval (CI) -0.90 to 0.22, [Analysis 1.1](#)).

1.2 Mental state: 2a. Average endpoint score of various scales (high = poor)

1.2.1 BPRS - anxiety/depression subscale score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 57) ([Boyer 1987](#)). We found evidence of a clear difference between 'fluphenazine (oral)' and 'amisulpride' within this subgroup (MD 2.60 95% CI 1.40 to 3.80, [Analysis 1.2](#)).

1.2.2 BPRS total score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 57) ([Boyer 1987](#)). There was not a clear difference between 'fluphenazine (oral)' and 'amisulpride' within this subgroup (MD 5.10 95% CI -2.35 to 12.55, [Analysis 1.2](#)).

1.2.3 SANS total score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 36) ([Saletu 1994](#)). There was a statistically significant difference (P = 0.03) favouring fluphenazine (oral) over amisulpride (MD -9.49 95% CI -17.88 to -1.10, [Analysis 1.2](#)).

1.3 Mental state: 2b. Average endpoint score of AMDP scale (high = poor)

1.3.1 short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing [Analysis 1.3](#).

1.4 Adverse effects: 1. Extrapyramidal side effects

1.4.1 concomitant anticholinergic medication - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 36) ([Saletu 1994](#)). There was a statistically significant difference (P = 0.04) favouring amisulpride over fluphenazine (oral) (risk ratio (RR) 7.82 95% CI 1.07 to 57.26, [Analysis 1.4](#)).

1.5 Leaving the study early

1.5.1 any reason

In this subgroup we found two trials (n = 98). There was no significant difference between the two treatment groups (RR 1.19 95% CI 0.63 to 2.28, [Analysis 1.5](#))

1.5.2 adverse effects - short term (up to 12 weeks)

In this subgroup we found two trials (n = 98). There was no significant difference between fluphenazine (oral) and amisulpride (RR 1.88 95% CI 0.24 to 14.68, [Analysis 1.5](#)).

1.5.1 inefficacy - short term (up to 12 weeks)

In this subgroup we found two trials (n = 98). There was no significant difference between fluphenazine (oral) and amisulpride (RR 1.82 95% CI 0.68 to 4.84, [Analysis 1.5](#)).

1.5.3 productive symptoms - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 36) ([Saletu 1994](#)). There was no significant difference between fluphenazine (oral) and amisulpride (RR 0.37 95% CI 0.04 to 3.25, [Analysis 1.5](#)).

COMPARISON 2: FLUPHENAZINE (ORAL) versus RISPERIDONE

2.1 Clinically important response (defined by study)

2.1.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 26) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.67 95% CI 0.13 to 3.35, Analysis 2.1).

2.2 Global state: 1. Average endpoint score of CGI scales (high = poor)

2.2.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 26) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (MD 0.07 95% CI -0.77 to 0.91, Analysis 2.2).

2.3 Mental state: 2a. Average endpoint scores (BPRS, high = poor)

2.3.1 BPRS endpoint total scale score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (MD -1.98 95% CI -12.96 to 9.00, Analysis 2.3).

2.3.2 BPRS positive subscale score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (MD -0.15 95% CI -4.22 to 3.92, Analysis 2.3).

2.3.3 BPRS negative subscale score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (MD -1.54 95% CI -3.94 to 0.86, Analysis 2.3).

2.3.4 BPRS hostile subscale score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (MD 0.92 95% CI -2.21 to 4.05, Analysis 2.3).

2.4 Mental state: 2b. Average endpoint score of various scales (high = poor) - short term (up to 12 weeks) (skewed data)

2.4.1 BPRS anxiety/depression subscale score

Data for this outcome are skewed and are best inspected by viewing Analysis 2.4.

2.4.2 BPRS activation subscale score

Data for this outcome are skewed and are best inspected by viewing Analysis 2.4.

2.5 Adverse effects: 1. Anticholinergic effect

2.5.1 blurred vision - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 1.08 95% CI 0.18 to 6.53, Analysis 2.5).

2.5.2 dry mouth - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 1.08 95% CI 0.18 to 6.53, Analysis 2.5).

2.5.3 urinary hesitancy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 5.38 95% CI 0.28 to 101.96, Analysis 2.5).

2.6 Adverse effects: 2. Central nervous system

2.6.1 anxiety - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.54 95% CI 0.06 to 5.24, Analysis 2.6).

2.6.2 headache - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.77 95% CI 0.33 to 1.79, Analysis 2.6).

2.6.3 lethargy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.81 95% CI 0.23 to 2.91, Analysis 2.6).

2.6.4 insomnia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 1.81 95% CI 0.55 to 5.98, Analysis 2.6).

2.6.5 somnolence - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.87 95% CI 0.30 to 2.49, Analysis 2.6).

2.7 Adverse effects: 3. Gastrointestinal

2.7.1 constipation - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 9.69 95% CI 0.58 to 163.02, Analysis 2.7).

2.7.2 diarrhoea - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.22 95% CI 0.01 to 4.08, Analysis 2.7).

2.7.3 dyspepsia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 3.25 95% CI 0.39 to 27.15, Analysis 2.7).

2.7.4 nausea/vomiting - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.72 95% CI 0.14 to 3.61, Analysis 2.7).

2.8 Adverse effects: 4. Other adverse events

2.8.1 urinary frequency - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.36 95% CI 0.02 to 8.05, Analysis 2.8).

2.8.2 orthostasis - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.43 95% CI 0.10 to 1.83, Analysis 2.8).

2.8.3 increased appetite - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.72 95% CI 0.14 to 3.61, Analysis 2.8).

2.8.4 dizziness - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and risperidone (RR 0.36 95% CI 0.04 to 3.02, Analysis 2.8).

2.9 Adverse effects: 5. Average endpoint weight loss (kg) (skewed data)

2.9.1 short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing Analysis 2.9

2.10 Leaving the study early

2.10.1 inefficacy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine

(oral) and risperidone (RR 1.08 95% CI 0.08 to 15.46, [Analysis 2.10](#)).

COMPARISON 3: FLUPHENAZINE (ORAL) versus QUETIAPINE

3.1 Clinically important response (defined by study)

3.1.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.62 95% CI 0.12 to 3.07, [Analysis 3.1](#)).

3.2 Global state: 1. CGI: Average endpoint CGI-SI (high = poor)

3.2.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (MD -0.03 95% CI -0.92 to 0.86, [Analysis 3.2](#)).

3.3 Mental state: 2a. General - average endpoint score (BPRS total, high = poor)

3.3.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (MD -1.98 95% CI -12.96 to 9.00, [Analysis 3.3](#)).

3.4 Mental state: 2b. Positive symptoms - average endpoint score (BPRS positive sub-score, high = poor)

3.4.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) ([Conley 2005](#)). There was a statistically significant difference (P = 0.0002) favouring fluphenazine (oral) over quetiapine (MD -13.61 95% CI -20.77 to -6.45, [Analysis 3.4](#)).

3.5 Mental state: 2c. Negative symptoms - average endpoint score (BPRS negative sub-score, high = poor)

3.5.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (MD -0.11 95% CI -2.27 to 2.05, [Analysis 3.5](#)).

3.6 Mental state: 2d. Anxiety/depression symptoms - average endpoint score (BPRS anxiety/depression sub-score, high score = poor)

3.6.1 short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing [Analysis 3.6](#).

3.7 Mental state: 2e. Hostility symptoms - average endpoint score (BPRS hostility sub-score, high = poor)

3.7.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (MD 0.79 95% CI -2.45 to 4.03, [Analysis 3.7](#)).

3.8 Mental state: 2f. Activation symptoms - average endpoint score (BPRS activation sub-score, high = poor)

3.8.1 short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing [Analysis 3.8](#).

3.9 Adverse effects: 1. Anticholinergic effect

3.9.1 dry mouth - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) ([Conley 2005](#)). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.17 to 5.98, [Analysis 3.9](#)).

3.9.2 blurred vision - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.17 to 5.98, Analysis 3.9).

3.9.3 urinary hesitancy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.17 to 5.98, Analysis 3.9).

3.10 Adverse effects: 2. Central nervous system

3.10.1 anxiety - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.07 to 14.21, Analysis 3.10).

3.10.2 headache - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.39 to 2.58, Analysis 3.10).

3.10.3 insomnia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.67 95% CI 0.51 to 5.46, Analysis 3.10).

3.10.4 lethargy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.50 95% CI 0.30 to 7.43, Analysis 3.10).

3.10.5 somnolence - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.33 95% CI 0.38 to 4.72, Analysis 3.10).

3.11 Adverse effects: 3. Gastrointestinal adverse effects

3.11.1 constipation - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.32 to 3.10, Analysis 3.11).

3.11.2 diarrhoea - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.20 95% CI 0.01 to 3.77, Analysis 3.11).

3.11.3 dyspepsia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 3.00 95% CI 0.36 to 24.92, Analysis 3.11).

3.11.4 nausea/ vomiting - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 2.00 95% CI 0.21 to 19.23, Analysis 3.11).

3.12 Adverse effects: 4a. Other adverse events

3.12.1 dizziness - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 1.00 95% CI 0.07 to 14.21, Analysis 3.12).

3.12.2 increased appetite - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.67 95% CI 0.13 to 3.30, Analysis 3.12).

3.12.3 orthostasis - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 2.00 95% CI 0.21 to 19.23, Analysis 3.12).

3.12.4 urinary frequency - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 24) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.33 95% CI 0.01 to 7.45, Analysis 3.12).

3.13 Adverse effects: 4b. Other - average endpoint weight loss (average weight in kg) (skewed)

3.13.1 short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing Analysis 3.13.

3.14 Leaving the study early

3.14.1 inefficacy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.46 95% CI 0.05 to 4.46, Analysis 3.14).

3.14.2 adverse effects - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 25) (Conley 2005). There was no significant difference between fluphenazine (oral) and quetiapine (RR 0.19 95% CI 0.01 to 3.52, Analysis 3.14).

COMPARISON 4: FLUPHENAZINE (ORAL) versus OLANZAPINE

4.1 Clinically important response (defined by author)

4.1.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 1.33 95% CI 0.86 to 2.07, Analysis 4.1).

4.1.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 1.60 95% CI 0.87 to 2.94, Analysis 4.1).

4.2 Global state: 1. CGI: Average change CGI-SI (high = poor)

4.2.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 0.70 95% CI -0.01 to 1.41, Analysis 4.2).

4.2.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was a statistically significant difference (P = 0.02) favouring olanzapine over fluphenazine (oral) (MD 0.90 95% CI 0.13 to 1.67, Analysis 4.2).

4.3 Mental state: 2a. General - average change score (BPRS total, high = poor)

4.3.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 7.10 95% CI -1.15 to 15.35, Analysis 4.3).

4.3.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 9.30 95% CI 0.10 to 18.50, Analysis 4.3).

4.4 Mental state: 2b. General - average change score (PANSS total, high = poor)

4.4.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 12.00 95% CI -2.03 to 26.03, Analysis 4.4).

4.4.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was a statistically significant difference (P = 0.04) favouring olanzapine over fluphenazine (oral) (MD 16.20 95% CI 0.41 to 31.99, Analysis 4.4).

4.5 Mental state: 2c. Positive symptoms - average change score (BPRS positive sub-score, high = poor)

4.5.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 2.30 95% CI -0.67 to 5.27, Analysis 4.5).

4.5.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 2.90 95% CI -0.29 to 6.09, Analysis 4.5).

4.6 Mental state: 2d. Positive symptoms - average endpoint score (PANSS positive sub-score, high = poor)

4.6.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was a statistically significant difference (P = 0.03) favouring fluphenazine (oral) over olanzapine (MD -5.10 95% CI -9.68 to -0.52, Analysis 4.6).

4.6.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was a statistically significant difference (P = 0.03) favouring fluphenazine (oral) over olanzapine (MD -5.10 95% CI -9.82 to -0.38, Analysis 4.6).

4.7 Mental state: 2e. Negative symptoms - average change score (BPRS negative sub-score, high = poor)

4.7.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 1.20 95% CI -0.47 to 2.87, Analysis 4.7).

4.7.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 55) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 1.70 95% CI -0.31 to 3.71, Analysis 4.7).

4.8 Mental state: 2f. Negative symptoms - average change score (PANSS negative sub-score, high = poor)

4.8.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 2.40 95% CI -0.96 to 5.76, Analysis 4.8).

4.8.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 3.00 95% CI -1.22 to 7.22, Analysis 4.8).

4.9 Mental state: 2g. General psychopathology - average change score (PANSS general psychopathology sub-score, high = poor)

4.9.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 6.20 95% CI -0.90 to 13.30, Analysis 4.9).

4.9.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was a statistically significant difference (P = 0.04) favouring olanzapine over fluphenazine (oral) (MD 8.20 95% CI 0.43 to 15.97, Analysis 4.9).

4.10 Mental state: 2h. Anxiety - average change score (HAMA, high = poor)

4.10.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 54) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 4.00 95% CI 0.08 to 7.92, Analysis 4.10).

4.10.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 59) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 6.00 95% CI -0.12 to 12.12, Analysis 4.10).

4.11 Satisfaction with treatment: 1. Average change score (DAI, low = poor)

4.11.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 52) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD -1.20 95% CI -2.44 to 0.04, Analysis 4.11).

4.11.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 52) (Dossenbach 1998). There was a statistically significant difference (P = 0.03) favouring olanzapine over between fluphenazine (oral) (MD -1.10 95% CI -2.08 to -0.12, Analysis 4.11).

4.12 Adverse effects: 1. General

4.12.1 at least one adverse effect - medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.04) favouring olanzapine over fluphenazine (oral) (RR 1.53 95% CI 1.02 to 2.31, Analysis 4.12).

4.13 Adverse effects: 2. Anticholinergic effect

4.13.1 concomitant anticholinergic medication - average endpoint dosage (mg/day) - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.001) favouring olanzapine over between fluphenazine (oral) (MD 0.89 95% CI 0.35 to 1.43, Analysis 4.13).

4.13.2 concomitant anticholinergic medication - average endpoint dosage (mg/day) - medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.008) favouring olanzapine over between fluphenazine (oral) (MD 1.08 95% CI 0.28 to 1.88, Analysis 4.13).

4.14 Adverse effects: 3a. Central nervous system

4.14.1 insomnia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 13.00 95% CI 0.76 to 220.96, Analysis 4.14).

4.15 Adverse effects: 3b. CNS (LSEQ, low = poor)

4.15.1 awakening from sleep average endpoint score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 53) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD -2.70 95% CI -10.18 to 4.78, Analysis 4.15).

4.15.2 behaviour following wakefulness average endpoint score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 53) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD -6.60 95% CI -13.92 to 0.72, Analysis 4.15).

4.15.3 getting to sleep average endpoint score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 53) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD -4.40 95% CI -14.18 to 5.38, Analysis 4.15).

4.15.4 sleep quality - average endpoint score - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 53) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD -6.10 95% CI -15.97 to 3.77, Analysis 4.15).

4.16 Adverse effects: 4a. Extrapyramidal effects

4.16.1 akathisia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 3.00 95% CI 0.90 to 10.01, Analysis 4.16).

4.16.2 hypertonia - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 3.00 95% CI 0.33 to 27.23, Analysis 4.16).

4.16.3 tremor - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 1.00 95% CI 0.22 to 4.56, Analysis 4.16).

4.17 Adverse effects: 4b. Extrapyramidal effects - average change score (SAS, high = poor)

4.17.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.001) favouring olanzapine over fluphenazine (oral) (MD 4.20 95% CI 1.68 to 6.72, Analysis 4.17).

4.17.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.008) favouring olanzapine over fluphenazine (oral) (MD 4.00 95% CI 1.02 to 6.98, Analysis 4.17).

4.18 Adverse effects: 4c. Extrapyramidal effects - average change score (HAS, high = poor)

4.18.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 59) (Dossenbach 1998). There was a statistically significant difference (P = 0.02) favouring olanzapine over fluphenazine (oral) (MD 6.60 95% CI 0.88 to 12.32, Analysis 4.18).

4.18.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 59) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 6.00 95% CI -0.12 to 12.12, Analysis 4.18).

4.19 Adverse effects: 4d. Extrapyramidal effects - average change score (AIMS, high = poor)

4.19.1 short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 1.10 95% CI -0.11 to 2.31, Analysis 4.19).

4.19.2 medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (MD 1.10 95% CI -0.45 to 2.65, Analysis 4.19).

4.20 Adverse effects: 5. Other adverse events

4.20.1 weight gain - medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 0.09 95% CI 0.01 to 1.57, Analysis 4.20).

4.21 Adverse effects: 5b. Other adverse events

4.21.1 concomitant anxiolytics medication average dosage (mg/day) - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.05) favouring olanzapine over fluphenazine (oral) (MD 4.65 95% CI 0.07 to 9.23, Analysis 4.21).

4.21.2 concomitant anxiolytics medication - average dosage (mg/day)- medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.03) favouring olanzapine over fluphenazine (oral) (MD 6.10 95% CI 0.63 to 11.57, Analysis 4.21).

4.21.3 effects on physiology - supine systolic blood pressure (average in mmHg) - short term (up to 12 weeks)

In this subgroup we only found one relevant trial (n = 60) (Dossenbach 1998). There was a statistically significant difference (P = 0.02) favouring fluphenazine (oral) over olanzapine (MD -10.00 95% CI -18.11 to -1.89, Analysis 4.21).

4.21.4 effects on physiology - supine systolic blood pressure (average in mmHg) - medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial ($n = 60$) (Dossenbach 1998). There was a statistically significant difference ($P = 0.02$) favouring fluphenazine (oral) over olanzapine (MD - 10.00 95% CI -18.11 to -1.89, Analysis 4.21).

4.22 Adverse effects: 5c. Other (skewed)

4.22.1 weight gain (average weight in kg) - short term (up to 12 weeks)

Data for this outcome are skewed and are best inspected by viewing Analysis 4.22.

4.22.2 weight gain (average weight in kg) - medium term (13 to 26 weeks)

Data for this outcome are skewed and are best inspected by viewing Analysis 4.22.

4.23 Leaving the study early

4.23.1 inefficacy - short term (up to 12 weeks)

In this subgroup we only found one relevant trial ($n = 60$) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 3.00 95% CI 0.33 to 27.23, Analysis 4.23).

4.23.2 adverse effects - medium term (13 to 26 weeks)

In this subgroup we only found one relevant trial ($n = 60$) (Dossenbach 1998). There was no significant difference between fluphenazine (oral) and olanzapine (RR 9.00 95% CI 0.51 to 160.17, Analysis 4.23).

5. SENSITIVITY ANALYSIS

5.1 Implication of randomisation

None of the included studies provided adequate details as to randomisation methods; furthermore, meta-analysis was not possible for our primary outcome, therefore the removal of studies with an inadequate description of randomisation left us with no data to compare.

5.2 Assumptions for lost binary data

Due to the relatively small loss to follow-up between studies, there was no difference in the estimate of effect of our primary outcome when we compared completer-only data with intention-to-treat analysis. Even when we assumed the extreme of each person leaving having a good outcome - this changed the findings by degree but not by direction and in no case changed the equivocal statistical significance of the results (Table 2).

5.3 Risk of bias

Each included study was rated as a 'high' risk of bias across at least one of the domains; again, meta-analysis was not possible for our primary outcome, therefore the removal of studies rated as a 'high' risk left us with no data to compare.

5.4 Imputed values

We did not include any cluster-randomised studies and therefore did not impute any ICC values.

5.5 Fixed-effect and random-effects

Since meta-analysis was not possible, there was no difference in the estimate of effect when using a fixed-effect model as opposed to a random-effects model.

6. Economic consideration of results

This review is one of several selected for economic consideration of findings. As yet, this has not been completed but should be available for next update.

ADDITIONAL SUMMARY OF FINDINGS *[Explanation]*

FLUPHENAZINE (ORAL) compared to RISPERIDONE for schizophrenia						
Patient or population: patients with schizophrenia Settings: USA Intervention: FLUPHENAZINE (ORAL) Comparison: RISPERIDONE						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	RISPERIDONE	FLUPHENAZINE (ORAL)				
Clinically important response (defined by study) - short term (up to 12 weeks) decreased rate of BPRS score < 20% Follow-up: 12 weeks	231 per 1000 ¹	155 per 1000 (30 to 773)	RR 0.67 (0.13 to 3.35)	26 (1 study)	⊕○○○ very low ^{2,3,4}	
Relapse (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Clinically important change in life skills (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Quality of life (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome

Adverse effects: Extrapyramidal effects - short/medium term (up to 12 weeks) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Leaving the study early - inefficacy short term (up to 12 weeks) Follow-up: 12 weeks	77 per 1000 ¹	83 per 1000 (6 to 1000)	RR 1.08 (0.08 to 15.46)	25 (1 study)	⊕○○○ very low ^{2,3,4}	
Cost-effectiveness (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome

*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 Control risk: mean baseline risk presented for single included study.

² Risk of bias: rated 'serious' - randomisation methods unclear; rated 'high' for attrition bias, with data for some included participants 'lost'; only one small study included (Conley 2005, n = 40).

³ Indirectness: rated 'serious' - only one included study provided data, which had three treatment arms (fluphenazine versus risperidone versus quetiapine).

⁴ Imprecision: rated 'very serious' - few participants, few events, leading to uncertainty in the precision of estimate of effect.

FLUPHENAZINE (ORAL) compared to QUETIAPINE for schizophrenia						
Patient or population: patients with schizophrenia Settings: USA Intervention: FLUPHENAZINE (ORAL) Comparison: QUETIAPINE						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	QUETIAPINE	FLUPHENAZINE (ORAL)				
Clinically important response (defined by study) - short term (up to 12 weeks) decreased rate of BPRS score < 20% Follow-up: 12 weeks	250 per 1000 ¹	155 per 1000 (30 to 767)	RR 0.62 (0.12 to 3.07)	25 (1 study)	⊕○○○ very low ^{2,3,4}	
Relapse (long term)	See comment	See comment	Not estimable	-	See comment	
Clinically important change in life skills (long term)	See comment	See comment	Not estimable	-	See comment	
Quality of life (long term)	See comment	See comment	Not estimable	-	See comment	
Adverse effects: Extrapyramidal effects - short/medium term (up to 12 weeks)	See comment	See comment	Not estimable	-	See comment	

Leaving the study early - inefficacy - short term (up to 12 weeks) Follow-up: 12 weeks	167 per 1000¹	77 per 1000 (8 to 743)	RR 0.46 (0.05 to 4.46)	25 (1 study)	⊕○○○ very low ^{2,3,4}
Cost-effectiveness (long term)	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 Control risk: mean baseline risk presented for single included study.

² Risk of bias: rated 'serious' - randomisation methods unclear; rated 'high' for attrition bias, with data for some included participants 'lost'; only one small study included (Conley 2005, n = 40).

³ Indirectness: rated 'serious' - only one included study provided data, which had three treatment arms (fluphenazine versus risperidone versus quetiapine).

⁴ Imprecision: rated 'very serious' - few participants, few events, leading to uncertainty in the precision of estimate of effect.

FLUPHENAZINE (ORAL) compared to OLANZAPINE for schizophrenia						
Patient or population: patients with schizophrenia Settings: multicentre, Croatia Intervention: FLUPHENAZINE (ORAL) Comparison: OLANZAPINE						
Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	OLANZAPINE	FLUPHENAZINE (ORAL)				
Clinically important response (defined by study) - short term (up to 12 weeks) decreased rate of PANSS score < 40%, decreased rate of BPRS score < 40% Follow-up: 22 weeks	500 per 1000 ¹	665 per 1000 (430 to 1000)	RR 1.33 (0.86 to 2.07)	60 (1 study)	⊕○○○ very low ^{2,3}	
Relapse (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Clinically important change in life skills (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome
Quality of life (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome

Adverse effects: Extrapyramidal effects - akathisia - short term (up to 12 weeks) Follow-up: 22 weeks	100 per 1000 ¹	300 per 1000 (90 to 1000)	RR 3.00 (0.90 to 10.01)	60 (1 study)	⊕○○○ very low ^{2,3}	
Leaving the study early: inefficacy - short term (up to 12 weeks) Follow-up: 22 weeks	33 per 1000 ¹	100 per 1000 (11 to 908)	RR 3.00 (0.33 to 27.23)	60 (1 study)	⊕○○○ very low ^{2,3}	
Cost-effectiveness (long term) - not reported	See comment	See comment	Not estimable	-	See comment	No study reported this outcome

*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 Control risk: mean baseline risk presented for single included study.

² Risk of bias: rated 'serious' - randomisation methods not adequately described; five participants excluded from analysis; sponsored by pharmaceutical company.

³ Imprecision: rated 'very serious' - few participants, few events, leading to uncertainty in the precision of estimate of effect (Dossenbach 1998, n = 60).

DISCUSSION

Summary of main results

1. FLUPHENAZINE (ORAL) versus AMISULPRIDE

1.1 Global state/mental state

Only one small study reported data the global state of participants in the short term (12 weeks). This showed no difference between fluphenazine and amisulpride. No trials provided longer-term data. No trials recorded relapse data. Another small study reported no clear difference between amisulpride and fluphenazine in the short term using the Brief Psychiatric Rating Scale (BPRS) (Analysis 1.2). The favourable outcome for negative symptoms reported by Saletu 1994 is, as always, of interest, but is only based on a short trial involving 36 people. We found no evidence that for these key outcomes there is any difference between fluphenazine and the newer drug - but all data are weak.

1.2 Adverse events

Less of the 19 allocated amisulpride in Saletu 1994 needed concomitant use of anticholinergic medication - a proxy measure of extrapyramidal symptoms. This would support most clinicians' experiences of fluphenazine having a high propensity to induce extrapyramidal side effects (EPSEs). There is significant concern in reporting bias with this small, drug company-funded study - and data on adverse events (the Webster Scale and Adverse Experience Scale) were not reported at all. Newer drugs are often marketed on having different and less problematic adverse effects than the old medications - so it is odd that there is not more confidence and openness in reporting.

1.3 Leaving the study early

One study found no significant difference for this outcome between the compounds.

2. FLUPHENAZINE (ORAL) versus RISPERIDONE

2.1 Global state/mental state

Only one study (n = 26) provided data on global state showing no difference between the compounds in the short term. There were no data on relapse. The same study did not show any significant difference between fluphenazine or risperidone on various scales of mental state response in the short term. This does seem to reflect the situation with amisulpride. No clear difference between the old and the new drug has been demonstrated.

2.2 Adverse effects

The same study found no difference between the compounds on a large number of individual adverse events. There is evidence that there is greater short-term weight gain with risperidone than fluphenazine. However, there is significant concern in the reporting of bias in this small, pharmaceutical company-funded study - data from the Simpson Angus Scale (SAS), the Barnes Akathisia Scale (BAS), and the Assessment of Involuntary Movements Scale (AIMS), Sexual Function: Changes in Sexual Function Questionnaire (CSFQ) and the Prolactin-Related Adverse Event Questionnaire (PRAEQ) were not reported. This is concerning as risperidone is associated with EPSEs and hyperprolactinaemia and, as with amisulpride, the reporting biases would tend to favour the newer drug.

2.3 Leaving the study early

The same study found no significant difference between the compounds for this outcome in the short term.

3. FLUPHENAZINE (ORAL) versus QUETIAPINE

3.1 Clinically important response (defined by study)/global state/mental state

Conley 2005 (n = 25) found no difference between the compounds in the short term in terms of 'clinically important response' and global state. For one measure on mental state, the same small short study found a difference favouring fluphenazine over quetiapine on the positive symptoms sub-score (Analysis 3.4). There were no demonstrated differences on the negative symptoms sub-score. So, in keeping with the other comparisons, tiny studies do not find convincing clinical differences between the old and the new drug.

3.2 Adverse effects

Again, as will the comparisons with other newer drugs, the same small study found no difference between the compounds on a large number of individual adverse events and reporting bias was considerable. If anything this reporting bias would have been favouring the newer and more expensive drug.

3.3 Leaving the study early

Conley 2005 found no difference between the compounds in the short term.

4. FLUPHENAZINE (ORAL) versus OLANZAPINE

4.1 Clinically important response (defined by study)/global state/mental state

Dossenbach 1998 (n = 60) found no difference between the compounds in the short term in terms of 'clinically important response'

in the short to mid term. The same study found no differences between compounds in the short term (average change on Cognitive Global Impression (CGI)), but a difference favoured olanzapine in the mid term (MD 0.9 95% CI 0.13 to 1.67, [Analysis 4.2](#)), though this is unlikely to be clinically significant. The same study found showed no clear differences between the compounds in the short term using BPRS or Positive and Negative Syndrome Scale (PANSS) overall. For other less important scores there was some favouring of olanzapine over fluphenazine and *vice versa*. The clinical significance of these findings is questionable due to small sample sizes and wide confidence intervals.

4.2 Adverse effects

Short- and mid-term data appeared to indicate less incidences of any adverse effects with olanzapine. Data support the expected outcome EPSEs with fluphenazine and marked short-term weight gain with olanzapine, but again, as with the other comparisons, there was the risk of reporting bias favouring the newer drug.

4.3 Leaving the study early

No difference was found between the compounds for attrition rates.

Overall completeness and applicability of evidence

1. Applicability

There were no international multicentre trials. The majority of patients were in-patients - thus these findings may not be applicable to the larger number of patients with schizophrenia now living in the community. Understandably, many of the exclusion criteria related to more severely ill patients (e.g. 'suicidality' or "acute paranoid psychosis") again bringing into question the applicability of these results in the acute in-patient setting. Similarly many other physical and mental co-morbidities such as addiction or depression were exclusion criteria - in reality such co-morbidities tend to be the norm rather than the exception.

Schizophrenia is a chronic, relapsing-remitting condition. None of the trials lasted longer than 22 weeks (and three of the four less than 12 weeks). Thus these trials cannot provide data on fluphenazine's role as a maintenance treatment in schizophrenia. Nor can they provide essential safety information about the long-term health implications of the studies drugs.

1.2 Quality of reporting

The four included studies failed to report several of our pre-specified secondary outcomes of interest, including economic outcomes, quality of life outcomes, service-use and hospitalisation

outcomes, relapse, and general function (such as social skills, employability). These are particularly patient-important outcomes that have been overlooked, and would add to the body of evidence regarding acceptability of treatment.

Each study had some examples of missing or unreported data due to attrition. Attrition is inevitable but unfortunately these studies had small-sample sizes. Only one gave details as to why the patients left the trial. Use of last observation carried forward (LOCF) was not clearly stated.

Other common tendencies affect clinically meaningful interpretation of the data - mean values were commonly reported without standard deviations. Most of the global state and mental state measurements were reported as continuous data that is difficult to interpret clinically. A more meaningful measure might be achieved by conversion to binary data such as "improved" or "not improved."

1.3 Heterogeneity

There was no heterogeneity between studies.

1.4 Publication bias

Formal tests to examine publication bias were underpowered. Therefore, we can not make a judgement in this regard.

Quality of the evidence

The quality of the findings were all rated as "very low." There were concerns with all four trials about a lack of detailed descriptions of the methodologies such as blinding and allocation practices. Most of the studies were pharmaceutical industry-funded. Not all outcomes were reported (particularly those related to adverse events such as akathisia scales), which was suggestive of reporting bias.

Potential biases in the review process

The review authors sought to adhere to the protocol, through the independent inspection of citations and full articles of potentially relevant studies.

Agreements and disagreements with other studies or reviews

We are not aware of other systematic reviews evaluating the effects of oral fluphenazine versus atypical antipsychotics in the treatment of schizophrenia.

AUTHORS' CONCLUSIONS

Implications for practice

1. For people with schizophrenia

These studies do not provide clear information about the relative merits or disadvantages of fluphenazine compared to the atypical antipsychotics many patients will be offered as first-line treatment. The data support the general point that use of fluphenazine carries the risk of many adverse effects including movement disorder. Many outcomes that patients will be concerned with such as tolerability and effect on quality of life of the drug are not answered by these studies.

2. For clinicians

These studies do not provide clear evidence to support or refute use of fluphenazine as first-line treatment for schizophrenia compared with atypical antipsychotics in terms of clinical response and effectiveness. As expected, the evidence suggests a greater propensity for EPSEs with fluphenazine than amisulpride or olanzapine, but more short-term weight gain with olanzapine and risperidone.

3. For managers or policy makers

Fluphenazine is inexpensive compared to atypical antipsychotics but there are no cost-effectiveness data. Likewise, there are no clear data relating to the relative effectiveness or patient satisfaction with the drug. It can cause significant side effects such as movement disorders.

Implications for research

1. General

Attempting to systematically review data on fluphenazine highlights the necessity of studies conforming to certain minimum criteria to allow extraction of clinically meaningful results. This would include more detailed and transparent study protocols giving full disclosure to such things as allocation, randomisation and blinding techniques. Use of more easily understandable binary outcomes could be helpful. It should also be the case that all data and results are fully reported. Two trials - [Ravanic 1996](#) (excluded study) and [Djukic-Dejanovic 2002](#) (awaiting assessment) are of direct relevance to this review but have no data that can be used.

We are not sure if the latter study should still be in awaiting assessment, or merged with [Ravanic 1996](#) as we continue to have no record of the full publication. Furthermore, excluded studies [Boyer 1986](#) and [Boyer 1996](#) may be one study. Close compliance with [CONSORT](#) would have helped clarify these issues.

2. Specific

2.1 Reviews

The excluded studies and the one awaiting assessment do pose important questions which would generate good comparisons for other reviews ([Table 3](#)).

2.2 Trials

It is difficult to derive meaningful clinical data to inform best practice with regard to the use of fluphenazine versus atypical antipsychotics in schizophrenia despite access to the pooled data of four different trials. Small sample sizes are problematic - it would be beneficial if international researchers were able to collaborate in more multicentre and long-term studies. Studies should also take into account more meaningful outcomes relating to hospital admissions, quality of life, mortality and cost-effectiveness. Given the current limitations in the literature, we propose a design for a new randomised trial ([Table 4](#)).

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We also thank Katie Jones for her help with data extraction of included studies.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Boyer 1987

Methods	Allocation: randomised. Blindness: not stated. Duration: 3 weeks washout plus 6 weeks treatment period. Setting: not stated. Design: parallel.
Participants	Diagnosis: schizophrenia (DSM-III). N = 62. Age: 21-53 years old. Sex: 43 men, 19 women. Duration ill: mean=12.3 years, SD=4.7 years. Inclusion criteria: duration ill between 1 to 20 years; absence of marked positive symptoms; score >7 on DSAS Exclusion criteria: not received antipsychotics in previous month
Interventions	1. Fluphenazine: 2 mg to 12 mg/day. N = 28. 2. Amisulpride: 50 mg to 300 mg/day. N = 34.
Outcomes	Mental state: BPRS.* Leaving the study early. Unable to use - Mental state: DSAS (not a validated scale). Behaviour: NOSIE (no SD reported). Adverse effects: physiological measures, CHESS (no data reported)
Notes	* The published papers clearly report SD as measure of variance but these seem to be SE and we treat them as such

Risk of bias

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "After a 3-week washout period, participants were randomly assigned" - no further details. (p.296)
Allocation concealment (selection bias)	Unclear risk	Not stated.
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Double-blind.

Boyer 1987 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not stated.
Incomplete outcome data (attrition bias) All outcomes	High risk	Five participants did not complete ratings for various BPRS components
Selective reporting (reporting bias)	High risk	Did not report complete data for NOSIE and CHES.
Other bias	Unclear risk	None obvious.

Conley 2005

Methods	Allocation: randomised, randomisation was performed by the dispensing pharmacy Blindness: double-blind. Duration: 12 weeks (with 4- to 6-week open-label qualification phase prior to randomisation) Setting: in-patients, Maryland (USA) Design: parallel.
Participants	Diagnosis: schizophrenia (DSM-III); therapy-refractory. N = 40. Age: 18-65 years old. Sex: men = 30 and women = 8. Length of illness: not stated. Included criteria: persistent positive psychotic symptoms at study entry ("moderate" severity ≥ 4 points on a 1- to 7-point scale) on 2 of 4 psychosis items on the BPRS scale; persistent global illness severity (BPRS total score ≥ 45 points on the 18-item scale and a CGI score of ≥ 4 points [moderately ill]); two prior failed treatment trials with two different antipsychotics at doses of 600 mg/day chlorpromazine equivalents, each of at least 6 weeks duration; no stable period of good social/occupational functioning within the previous 5 years Exclusion criteria: not stated.
Interventions	1. Fluphenazine: mean, 13.2 mg/day, SD 1.17 mg/day, n = 13. 2. Risperidone: mean, 4.31 mg/day, SD 0.63 mg/day, n = 13. 3. Quetiapine: mean, 463.6 mg/day, SD 50.5 mg/day, n = 12.
Outcomes	Clinically important response: no clinical improvement*. Global state: CGI severity score. Mental state: BPRS (global score; negative, positive, anxiety-depression score, hostility, activation score) Adverse effects. Leave the study early. Unable to use - Simpson Angus Scale (SAS), the Barnes Akathisia Scale (BAS), and the Assessment of Involuntary Movements Scale (AIMS), Sexual Function: Changes in Sexual Function

Conley 2005 (Continued)

	Questionnaire (CSFQ) (no data reported) Quality of life (no SD reported). Adverse effect: the Prolactin-Related Adverse Event Questionnaire (PRAEQ) (no data reported)
Notes	Funding: National Institutes of Mental Health (NIMH grant MH-47311); study medications supplied by Janssen Pharmaceutica and Astra-Zeneca Pharmaceuticals * decreased rate of BPRS score < 20%.

<i>Risk of bias</i>		<i>Risk of bias</i>
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomisation was performed by the dispensing pharmacy".(p.341)
Allocation concealment (selection bias)	Unclear risk	Not stated.
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "After a 4-6 week open-label trial with either olanzapine (or a typical antipsychotic other than fluphenazine), participants who did not achieve a 20% reduction in their total BPRS scores and who still had a total BPRS ≥ 35 points were randomly assigned. After open-label phase, participants were randomised to double blind study." No further details.(p.164)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not stated.
Incomplete outcome data (attrition bias) All outcomes	High risk	N = 40 participants randomised - n = 2 pieces of data "lost" (p.165); only results for 38 participants reported
Selective reporting (reporting bias)	High risk	SAS, BAS, AIMS, NOSIE, CHES, CSFQ, PRAEQ were not well-reported
Other bias	Unclear risk	Funding: National Institutes of Mental Health (NIMH grant MH-47311), study medications supplied by Janssen Pharmaceutica and Astra-Zeneca Pharmaceuticals

Dossenbach 1998

Methods	<p>Allocation: randomised, no further information.</p> <p>Blindness: double-blind.</p> <p>Duration: Acute phase: 6 weeks (2 to 9 days placebo lead-in); long term: 22 weeks</p> <p>Setting: hospital, multicentre, Croatia.</p> <p>Design: parallel.</p>
Participants	<p>Diagnosis: schizophrenia or schizoaffective disorder (DSM-IV)</p> <p>N = 60.</p> <p>Age: mean-35.4 years, SD-10.4 years.</p> <p>Sex: men = 28 and women = 32.</p> <p>Length of illness: not stated.</p> <p>Inclusion criteria: schizophrenia, BPRS \geq 42, CGI \geq 4 .</p> <p>Exclusion criteria: pregnant or lactating women, serious or unstable illness; history of intolerance to olanzapine; DSM substance dependence excluding caffeine or nicotine within last 30 days; serious suicide risk; significantly elevated liver function results; active hepatitis B or current jaundice; received treatment with injectable neuroleptic within less than one dosing interval between depot neuroleptic injection prior to study entry; previously intolerant or non-responsive to fluphenazine; previous participation in any olanzapine clinical trial; pregnancy or lactating; uncorrected hypothyroidism or hyperthyroidism, myasthenia gravis, narrow-angle glaucoma, chronic urinary retention and/or clinically significant prostatic hypertrophy, a history of seizures, severe allergies or multiple adverse drug reactions, a history of leukopenia without known aetiology</p>
Interventions	<p>1. Fluphenazine: 5 mg to 20 mg/day, average, 11.7 \pm 3.0 mg/day for acute phase (6 weeks) and 11.7 \pm 3.0 mg/day for long term (22 weeks), n = 30</p> <p>2. Olanzapine: 6 mg to 21 mg/day, average, 13.6 \pm 2.4 mg/day for acute phase (6 weeks) and 14.8 \pm 2.5 mg/day for long term (22 weeks), n = 30</p>
Outcomes	<p>Clinically important response: no clinical improvement*.</p> <p>Global state: CGI severity change score.</p> <p>Mental state: BPRS (global score; negative, positive change score); PANSS (total, positive, negative, general psychopathology change score), Hillside Akathisia Scale (HAS)</p> <p>Quality of Sleep scale: Leeds Sleep Evaluation Questionnaire (LSEQ) total and subscale score</p> <p>Satisfaction: Drug Attitude Inventory (DAI).</p> <p>Extrapyramidal adverse effects: Simpson Angus Scale (SAS), and the Assessment of Involuntary Movements Scale (AIMS)</p> <p>Leave the study early.</p> <p>Unable to use: vital signs, ECG, laboratory findings, no data reported. HAMA subscale score. LSEQ in medium term, data not reported</p>
Notes	<p>Funding: Eli Lilly and Company.</p> <p>*Two definitions: decreased rate of PANSS score < 40%, decreased rate of BPRS score < 40%. The data were reported separately in our data analysis based on these two definitions</p>

Risk of bias
Risk of bias

Bias	Authors' judgement	Support for judgement
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Dossenbach 1998 (Continued)

Random sequence generation (selection bias)	Unclear risk	Quotes: “Random allocation at a 1:1 ratio” (p.312)
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: “This was a long-term, randomised, double-blind parallel clinical trial” (p.312)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: “This was a long-term, randomised, double-blind parallel clinical trial” (p.312)
Incomplete outcome data (attrition bias) All outcomes	High risk	Out of 60 participants, five (n = 3 olanzapine; n = 2 fluphenazine) were excluded from efficacy analysis because they did not meet inclusion criteria for BPRS or CGI. Three participants missing from DAI results because of no baseline data for two (n = 1 olanzapine; n = 1 fluphenazine) and one participant on fluphenazine discontinuing without having a DAI performed. Four participants in fluphenazine group discontinued because of adverse event. All participants were included in safety analysis
Selective reporting (reporting bias)	Low risk	All measured outcomes were well reported.
Other bias	High risk	Funding: sponsored by Eli Lilly and Company.

Saletu 1994

Methods	Allocation: randomised, no further details. Blindness: double-blind. Duration: 3-day washout period plus 6 weeks treatment period Setting: in-patients, Austria. Design: parallel.
Participants	Diagnosis: DSM-III schizophrenia. N = 40. Age: mean-31 years, SD-6.4 years. Sex: men = 32 and women = 8. Length of illness: mean 82.6-118.3 months. Inclusion criteria: clinical diagnosis (ICD 9 criteria) of either a simple type (295.0), hebephrenic type (295.1) or residual type (295.6) of schizophrenia; minimal age of 18 years; minimal length of illness of 1 year; a necessity of 6 weeks in-patient treatment Exclusion criteria: an acute phase of a paranoid schizophrenia; pronounced symptoms of

	depression, neurotic asthenia or neurotic depression; reactive depressive psychosis; alcohol-induced psychiatric disturbances; gravidity; physical illness; treatment with lithium salts; treatment with depot neuroleptics within the last 45 days; potential premature discontinuation of treatment
Interventions	1. Fluphenazine: 2 mg/day to 4 mg/day, n = 21. 2. Amisulpride: 50 mg/day to 100 mg/day, n = 19. During the first 2 weeks, the dosage consisted of single doses of 50 mg/day amisulpride or 2 mg/day fluphenazine. From the third week up to the sixth, the daily doses were 100 mg amisulpride (50 mg twice daily) and 4 mg fluphenazine (2 mg twice daily)
Outcomes	Global state: CGI Mental state: Association for Methodology and Documentation in Psychiatry (AMDP), SANS Extrapyramidal adverse effects: Webster scale, contaminant of anticholinergic drugs Adverse effect: Adverse experience questionnaire. Leave the study early. Unable to use: Global function: Grunberger AD test (Alphabetical cross-out test); Grunberger psychomotor activity test; numerical memory test; Pauli test; reaction time test; complex reaction test; Zerssen well-being scale; semantic differential polarity profile; state-trait anxiety scale; CFF descending threshold; skin conductance (mean); skin conductance fluctuations. There is no SD reported and no reply from the author Webster scale, Adverse experience questionnaire, EEG, no data reported on this outcome
Notes	Funding: Experimental drugs supplied by Synthelabo Recherche/Laboratoires Delagrangre (Bagneux, France)

<i>Risk of bias</i>		<i>Risk of bias</i>
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "patients were included in the double-blind, parallel group study" (p.127)
Allocation concealment (selection bias)	Unclear risk	Allocation concealment not described.
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Quote: "patients were included in the double-blind, parallel group study" (p.127)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "patients were included in the double-blind, parallel group study" (p.127)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Out of 40 participants, five discontinued therapy prematurely. Three amisulpride participants dropped out due to productive symptoms (days 14, 28 and 35) while two

Saletu 1994 (Continued)

		fluphenazine participants dropped out due to depressive symptoms (days 21 and 28)
Selective reporting (reporting bias)	High risk	Webster scale, adverse experience questionnaire and EEG were not reported
Other bias	High risk	Funding: Experimental drugs supplied by Synthelabo Recherche/Laboratoires Delagrangre (Bagneux, France)

AIMS - Assessment of Involuntary Movements Scale
 AMDP - Association for Methodology and Documentation in Psychiatry
 BAS - Barnes Akathisia Scale
 BPRS - Brief Psychiatric Rating Scale
 CFF: critical flicker frequency
 CGI - Cognitive Global Impression
 CHES - Changes in Health, End-Stage, Disease, Signs, and Symptoms
 CSFQ - Changes in Sexual Functioning Questionnaire
 DAI - Drug Attitude Inventory
 DSAS - Depression Anxiety and Stress Scale
 DSM-III: Diagnostic and Statistical Manual, third edition
 DSM-IV: Diagnostic and Statistical Manual, fourth edition
 ECG - electrocardiogram
 HAMA - Hamilton Anxiety Scale
 HAS - Hillside Akathisia Scale
 ICD: International Classification of Diseases
 LSEQ - Leeds Sleep Evaluation Questionnaire
 NOSIE - Nurses' Observation Scale for Inpatient Evaluation
 PANSS - Positive and Negative Syndrome Scale
 PRAEQ - Prolactin Related Adverse Event Questionnaire
 SANS - Scale for Assessment of Negative Symptoms
 SAS - Simpson-Angus Scale
 SD: standard deviation
 SE: standard error

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Boyer 1986	Allocation: randomised. Participants: people with schizophrenia. Interventions: amisulpride versus haloperidol (n = 39), not fluphenazine We are unclear if this study is one report of a larger multicentre trial (Boyer 1996).

(Continued)

Boyer 1987a	Allocation: randomised. Participants: people with schizophrenia. Interventions: amisulpride 100 mg versus amisulpride 300 mg versus placebo, not fluphenazine
Boyer 1996	Allocation: randomised. Participants: people with schizophrenia. Interventions: amisulpride versus haloperidol (n = 191), not fluphenazine We are unclear if this study is the final report of trial also reported by individual centre (Boyer 1986).
Pickar 1992	Allocation: not randomised, one arm cross-over design.
Ravanic 1996	Allocation: randomised. Participants: schizophrenia DSM III-R. Intervention: fluphenazine versus clozapine. Outcomes: no usable data.

DSM-III-R: Diagnostic and Statistical Manual, third edition, revised

Characteristics of studies awaiting assessment *[ordered by study ID]*

Djukic-Dejanovic 2002

Methods	Allocation: randomised.
Participants	Diagnosis: schizophrenia (DSM-IV). N = 44.
Interventions	1. Fluphenazine. N = 10. 2. Clozapine. N = 23. 3. Haloperidol. N = 11.
Outcomes	Adverse effects.
Notes	Full paper required (conference abstract only) We are very unclear if this study is one further report of the Ravanic 1996 trial. Numbers of participants, and the short description is identical but interventions are similar but not the same

DSM-IV: Diagnostic and Statistical Manual, fourth edition

DATA AND ANALYSES

Comparison 1. FLUPHENAZINE (ORAL) vs AMISULPRIDE

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Global state: 1. Average endpoint score of CGI scales (high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 short term (up to 12 weeks)	1	36	Mean Difference (IV, Random, 95% CI)	-0.34 [-0.90, 0.22]
2 Mental state: 2a. Average endpoint score of various scales (high = poor)	2		Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 BPRS - anxiety/depression subscale score - short term (up to 12 weeks)	1	57	Mean Difference (IV, Random, 95% CI)	2.60 [1.40, 3.80]
2.2 BPRS total score - short term (up to 12 weeks)	1	57	Mean Difference (IV, Random, 95% CI)	5.10 [-2.35, 12.55]
2.3 SANS total score - short term (up to 12 weeks)	1	36	Mean Difference (IV, Random, 95% CI)	-9.49 [-17.88, -1.10]
3 Mental state: 2b. Average endpoint score of AMDP scale (high = poor)			Other data	No numeric data
3.1 short term (up to 12 weeks)			Other data	No numeric data
4 Adverse effects: 1. Extrapyramidal effects	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
4.1 concomitant anticholinergic medication - short term (up to 12 weeks)	1	36	Risk Ratio (M-H, Random, 95% CI)	7.82 [1.07, 57.26]
5 Leaving the study early	2		Risk Ratio (IV, Random, 95% CI)	Subtotals only
5.1 any reason	2	98	Risk Ratio (IV, Random, 95% CI)	1.19 [0.63, 2.28]
5.2 adverse effects - short term (up to 12 weeks)	2	98	Risk Ratio (IV, Random, 95% CI)	1.88 [0.24, 14.68]
5.3 inefficacy - short term (up to 12 weeks)	2	98	Risk Ratio (IV, Random, 95% CI)	1.82 [0.68, 4.84]
5.4 productive symptoms - short term (up to 12 weeks)	1	36	Risk Ratio (IV, Random, 95% CI)	0.37 [0.04, 3.25]

Comparison 2. FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Clinically important response (defined by study)	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
1.1 short term (up to 12 weeks)	1	26	Risk Ratio (M-H, Random, 95% CI)	0.67 [0.13, 3.35]
2 Global state: 1. Average endpoint score of CGI scales (high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 short term (up to 12 weeks)	1	26	Mean Difference (IV, Random, 95% CI)	0.07 [-0.77, 0.91]
3 Mental state: 2a. Average endpoint scores (BPRS, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 BPRS endpoint total scale score - short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-1.98 [-12.96, 9.00]
3.2 BPRS positive subscale score - short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-0.15 [-4.22, 3.92]
3.3 BPRS negative subscale score - short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-1.54 [-3.94, 0.86]
3.4 BPRS hostile subscale score - short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	0.92 [-2.21, 4.05]
4 Mental state: 2b. Average endpoint score of various scales (high = poor) - short term (up to 12 weeks) (skewed data)			Other data	No numeric data
4.1 BPRS anxiety/depression subscale score			Other data	No numeric data
4.2 BPRS activation subscale score			Other data	No numeric data
5 Adverse effects: 1. Anticholinergic effect	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
5.1 blurred vision - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	1.08 [0.18, 6.53]
5.2 dry mouth - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	1.08 [0.18, 6.53]
5.3 urinary hesitancy - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	5.38 [0.28, 101.96]
6 Adverse effects: 2. Central nervous system	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
6.1 anxiety - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.54 [0.06, 5.24]

6.2 headache - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.77 [0.33, 1.79]
6.3 lethargy - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.81 [0.23, 2.91]
6.4 insomnia - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	1.81 [0.55, 5.98]
6.5 somnolence - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.87 [0.30, 2.49]
7 Adverse effects: 3. Gastrointestinal	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
7.1 constipation - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	9.69 [0.58, 163.02]
7.2 diarrhoea - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.22 [0.01, 4.08]
7.3 dyspepsia - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	3.25 [0.39, 27.15]
7.4 nausea/vomiting - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.72 [0.14, 3.61]
8 Adverse effects: 4. Other adverse events	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
8.1 urinary frequency - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.36 [0.02, 8.05]
8.2 orthostasis - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.43 [0.10, 1.83]
8.3 increased appetite - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.72 [0.14, 3.61]
8.4 dizziness - short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.36 [0.04, 3.02]
9 Adverse effects: 5. Average endpoint weight loss (kg) (skewed data)			Other data	No numeric data
9.1 short term (up to 12 weeks)			Other data	No numeric data
10 Leaving the study early	1		Risk Ratio (IV, Random, 95% CI)	Subtotals only
10.1 inefficacy - short term (up to 12 weeks)	1	25	Risk Ratio (IV, Random, 95% CI)	1.08 [0.08, 15.46]

Comparison 3. FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Clinically important response (defined by study)	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
1.1 short term (up to 12 weeks)	1	25	Risk Ratio (M-H, Random, 95% CI)	0.62 [0.12, 3.07]
2 Global state: 1. CGI: Average endpoint CGI-SI (high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only

2.1 short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-0.03 [-0.92, 0.86]
3 Mental state: 2a. General - average endpoint score (BPRS total, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-1.98 [-12.96, 9.00]
4 Mental state: 2b. Positive symptoms - average endpoint score (BPRS positive sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 short term (up to 12 weeks)	1	25	Mean Difference (IV, Random, 95% CI)	-13.61 [-20.77, -6.45]
5 Mental state: 2c. Negative symptoms - average endpoint score (BPRS negative sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 short term (up to 12 weeks)	1	24	Mean Difference (IV, Random, 95% CI)	-0.11 [-2.27, 2.05]
6 Mental state: 2d. Anxiety/depression symptoms - average endpoint score (BPRS anxiety/depression sub-score, high score = poor)			Other data	No numeric data
6.1 short term (up to 12 weeks)			Other data	No numeric data
7 Mental state: 2e. Hostility symptoms - average endpoint score (BPRS hostility sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
7.1 short term (up to 12 weeks)	1	24	Mean Difference (IV, Random, 95% CI)	0.79 [-2.45, 4.03]
8 Mental state: 2f. Activation symptoms - average endpoint score (BPRS activation sub-score, high = poor)			Other data	No numeric data
8.1 short term (up to 12 weeks)			Other data	No numeric data
9 Adverse effects: 1. Anticholinergic effect	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
9.1 dry mouth - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.17, 5.98]
9.2 blurred vision - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.17, 5.98]
9.3 urinary hesitancy - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.17, 5.98]
10 Adverse effects: 2. Central nervous system	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
10.1 anxiety - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.07, 14.21]

10.2 headache - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.39, 2.58]
10.3 insomnia - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.67 [0.51, 5.46]
10.4 lethargy - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.5 [0.30, 7.43]
10.5 somnolence - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.33 [0.38, 4.72]
11 Adverse effects: 3. Gastrointestinal adverse effects	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
11.1 constipation - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.32, 3.10]
11.2 diarrhoea - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	0.2 [0.01, 3.77]
11.3 dyspepsia - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	3.0 [0.36, 24.92]
11.4 nausea/ vomiting - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	2.0 [0.21, 19.23]
12 Adverse effects: 4a. Other adverse events	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
12.1 dizziness - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	1.0 [0.07, 14.21]
12.2 increased appetite - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	0.67 [0.13, 3.30]
12.3 orthostasis - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	2.0 [0.21, 19.23]
12.4 urinary frequency - short term (up to 12 weeks)	1	24	Risk Ratio (M-H, Random, 95% CI)	0.33 [0.01, 7.45]
13 Adverse effects: 4b. Other - average endpoint weight loss (average weight in kg) (skewed)			Other data	No numeric data
13.2 short term (up to 12 weeks)			Other data	No numeric data
14 Leaving the study early	1		Risk Ratio (IV, Random, 95% CI)	Subtotals only
14.1 inefficacy - short term (up to 12 weeks)	1	25	Risk Ratio (IV, Random, 95% CI)	0.46 [0.05, 4.46]
14.2 adverse effects - short term (up to 12 weeks)	1	25	Risk Ratio (IV, Random, 95% CI)	0.19 [0.01, 3.52]

Comparison 4. FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Clinically important response (defined by author)	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
1.1 short term (up to 12 weeks)	1	60	Risk Ratio (M-H, Random, 95% CI)	1.33 [0.86, 2.07]

1.2 medium term (13 to 26 weeks)	1	60	Risk Ratio (M-H, Random, 95% CI)	1.6 [0.87, 2.94]
2 Global state: 1. CGI: Average change CGI-SI (high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 short term (up to 12 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	0.7 [-0.01, 1.41]
2.2 medium term (13 to 26 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	0.90 [0.13, 1.67]
3 Mental state: 2a. General - average change score (BPRS total, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 short term (up to 12 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	7.1 [-1.15, 15.35]
3.2 medium term (13 to 26 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	9.3 [0.10, 18.50]
4 Mental state: 2b. General - average change score (PANSS total, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 short term (up to 12 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	12.0 [-2.03, 26.03]
4.2 medium term (13 to 26 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	16.20 [0.41, 31.99]
5 Mental state: 2c. Positive symptoms - average change score (BPRS positive sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 short term (up to 12 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	2.3 [-0.67, 5.27]
5.2 medium term (13 to 26 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	2.90 [-0.29, 6.09]
6 Mental state: 2d. Positive symptoms - average endpoint score (PANSS positive sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
6.1 short term (up to 12 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	-5.1 [-9.68, -0.52]
6.2 medium term (13 to 26 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	-5.1 [-9.82, -0.38]
7 Mental state: 2e. Negative symptoms - average change score (BPRS negative sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
7.1 short term (up to 12 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	1.20 [-0.47, 2.87]
7.2 medium term (13 to 26 weeks)	1	55	Mean Difference (IV, Random, 95% CI)	1.70 [-0.31, 3.71]
8 Mental state: 2f. Negative symptoms - average change score (PANSS negative sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only

8.1 short term (up to 12 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	2.40 [-0.96, 5.76]
8.2 medium term (13 to 26 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	3.0 [-1.22, 7.22]
9 Mental state: 2g. General psychopathology - average change score (PANSS general psychopathology sub-score, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
9.1 short term (up to 12 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	6.20 [-0.90, 13.30]
9.2 medium term (13 to 26 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	8.2 [0.43, 15.97]
10 Mental state: 2h. Anxiety - average change score (HAMA, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
10.1 short term (up to 12 weeks)	1	54	Mean Difference (IV, Random, 95% CI)	4.00 [0.08, 7.92]
10.2 medium term (13 to 26 weeks)	1	59	Mean Difference (IV, Random, 95% CI)	6.0 [-0.12, 12.12]
11 Satisfaction with treatment: 1. Average change score (DAI, low = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
11.1 short term (up to 12 weeks)	1	52	Mean Difference (IV, Random, 95% CI)	-1.2 [-2.44, 0.04]
11.2 medium term (13 to 26 weeks)	1	52	Mean Difference (IV, Random, 95% CI)	-1.1 [-2.08, -0.12]
12 Adverse effects: 1. General	1		Risk Ratio (IV, Random, 95% CI)	Subtotals only
12.1 at least one adverse effect - medium term (13 to 26 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	1.53 [1.02, 2.31]
13 Adverse effects: 2. Anticholinergic effect	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
13.1 concomitant anticholinergic medication - average endpoint dosage (mg/d) - short term (up to 12 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	0.89 [0.35, 1.43]
13.2 concomitant anticholinergic medication - average endpoint dosage (mg/d) - medium term (13 to 26 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	1.08 [0.28, 1.88]
14 Adverse effects: 3a. Central nervous system	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
14.1 insomnia - short term (up to 12 weeks)	1	60	Risk Ratio (M-H, Random, 95% CI)	13.0 [0.76, 220.96]
15 Adverse effects: 3b. CNS (LSEQ, low = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only

15.1 awakening from sleep average endpoint score - short term (up to 12 weeks)	1	53	Mean Difference (IV, Random, 95% CI)	-2.7 [-10.18, 4.78]
15.2 behaviour following wakefulness average endpoint score - short term (up to 12 weeks)	1	53	Mean Difference (IV, Random, 95% CI)	-6.60 [-13.92, 0.72]
15.3 getting to sleep average endpoint score - short term (up to 12 weeks)	1	53	Mean Difference (IV, Random, 95% CI)	-4.4 [-14.18, 5.38]
15.4 sleep quality - average endpoint score - short term (up to 12 weeks)	1	53	Mean Difference (IV, Random, 95% CI)	-6.10 [-15.97, 3.77]
16 Adverse effects: 4a. Extrapyramidal effects	1		Risk Ratio (IV, Random, 95% CI)	Subtotals only
16.1 akathisia - short term (up to 12 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	3.0 [0.90, 10.01]
16.2 hypertonia - short term (up to 12 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	3.0 [0.33, 27.23]
16.3 tremor - short term (up to 12 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	1.0 [0.22, 4.56]
17 Adverse effects: 4b. Extrapyramidal effects - average change score (SAS, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
17.1 short term (up to 12 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	4.20 [1.68, 6.72]
17.2 medium term (13 to 26 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	4.0 [1.02, 6.98]
18 Adverse effects: 4c. Extrapyramidal effects - average change score (HAS, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
18.1 short term (up to 12 weeks)	1	59	Mean Difference (IV, Random, 95% CI)	6.6 [0.88, 12.32]
18.2 medium term (13 to 26 weeks)	1	59	Mean Difference (IV, Random, 95% CI)	6.0 [-0.12, 12.12]
19 Adverse effects: 4d. Extrapyramidal effects - average change score (AIMS, high = poor)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
19.1 short term (up to 12 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	1.1 [-0.11, 2.31]
19.2 medium term (13 to 26 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	1.1 [-0.45, 2.65]
20 Adverse effects: 5. Other adverse events	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
20.1 weight gain - medium term (13 to 26 weeks)	1	60	Risk Ratio (M-H, Random, 95% CI)	0.09 [0.01, 1.57]
21 Adverse effects: 5b. Other adverse events	1		Mean Difference (IV, Random, 95% CI)	Subtotals only

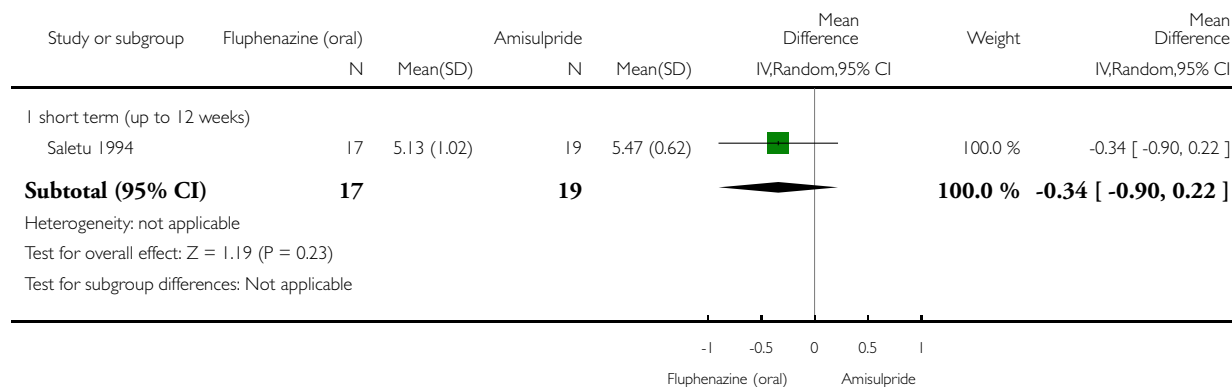
21.1 concomitant anxiolytics medication average dosage (mg/d) - short term (up to 12 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	4.65 [0.07, 9.23]
21.2 concomitant anxiolytics medication - average dosage (mg/d)- medium term (13 to 26 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	6.10 [0.63, 11.57]
21.3 effects on physiology - supine systolic blood pressure (average in mmHg) - short term (up to 12 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	-10.0 [-18.11, -1.89]
21.4 effects on physiology - supine systolic blood pressure (average in mmHg) - medium term (13 to 26 weeks)	1	60	Mean Difference (IV, Random, 95% CI)	-10.0 [-18.11, -1.89]
22 Adverse effects: 5c. Other (skewed)			Other data	No numeric data
22.1 weight gain (average weight in kg) - short term (up to 12 weeks)			Other data	No numeric data
22.2 weight gain (average weight in kg) - medium term (13 to 26 weeks)			Other data	No numeric data
23 Leaving the study early	1		Risk Ratio (IV, Random, 95% CI)	Subtotals only
23.1 inefficacy - short term (up to 12 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	3.0 [0.33, 27.23]
23.2 adverse effects - medium term (13 to 26 weeks)	1	60	Risk Ratio (IV, Random, 95% CI)	9.00 [0.51, 160.17]

Analysis 1.1. Comparison 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE, Outcome 1 Global state: 1. Average endpoint score of CGI scales (high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE

Outcome: 1 Global state: 1. Average endpoint score of CGI scales (high = poor)

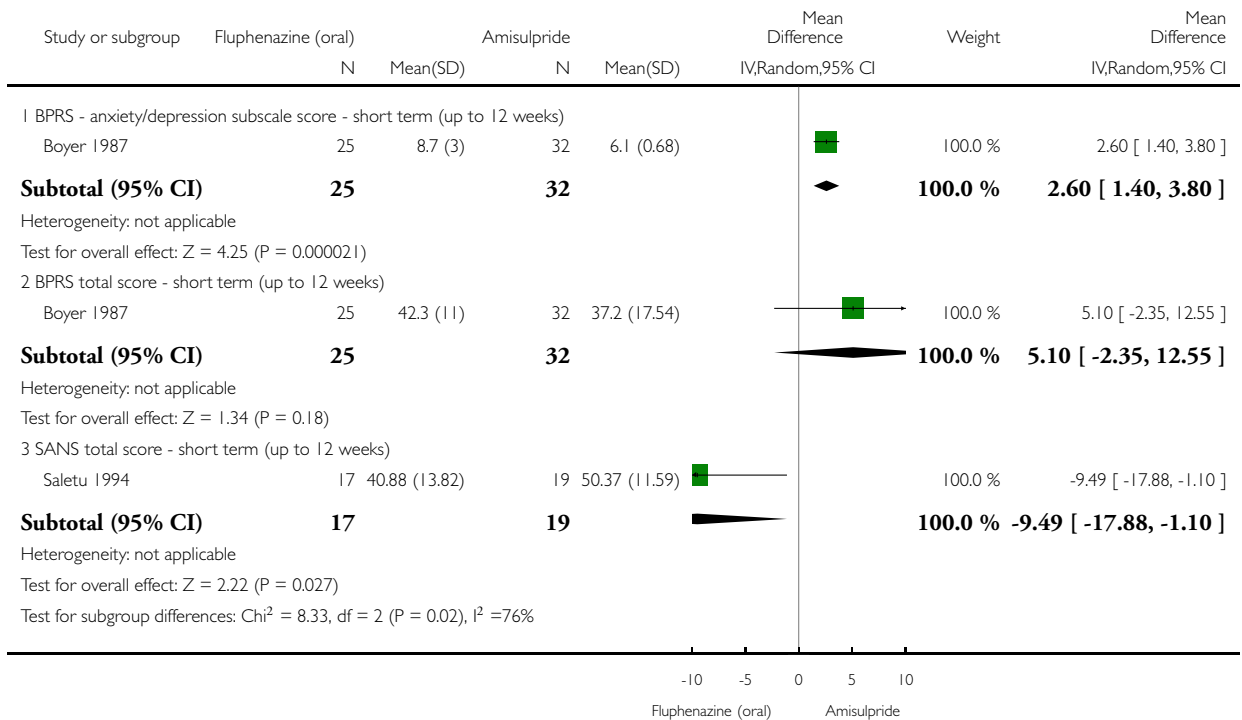


Analysis 1.2. Comparison 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE, Outcome 2 Mental state: 2a. Average endpoint score of various scales (high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE

Outcome: 2 Mental state: 2a. Average endpoint score of various scales (high = poor)



Analysis 1.3. Comparison 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE, Outcome 3 Mental state: 2b. Average endpoint score of AMDP scale (high = poor).

Mental state: 2b. Average endpoint score of AMDP scale (high = poor)

Study	Interventions	Mean	SD	N
short term (up to 12 weeks)				

Mental state: 2b. Average endpoint score of AMDP scale (high = poor) (Continued)

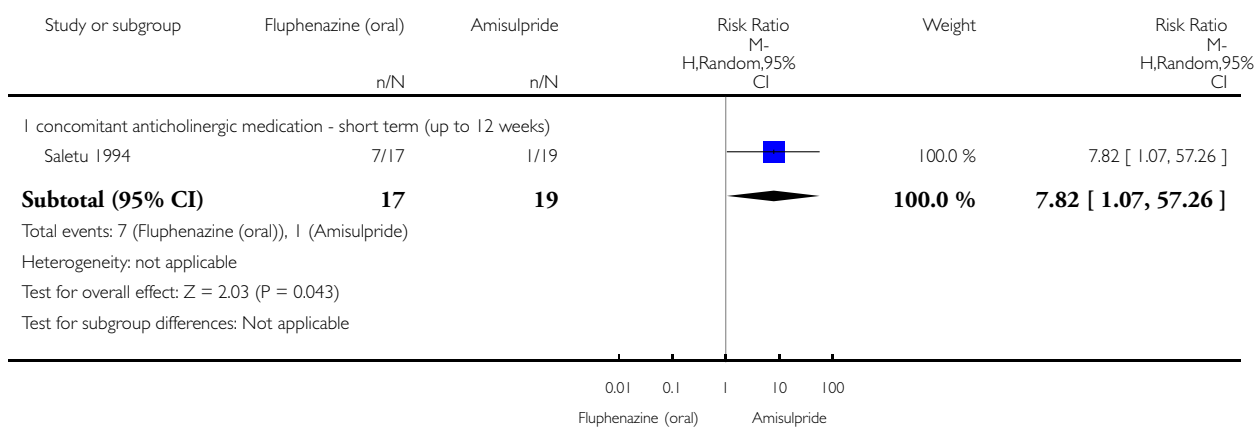
Saletu 1994	Fluphenazine	7.13	4.26	17
Saletu 1994	Amisulpride	8.76	4.13	19

Analysis 1.4. Comparison 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE, Outcome 4 Adverse effects: 1. Extrapyramidal effects.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE

Outcome: 4 Adverse effects: 1. Extrapyramidal effects

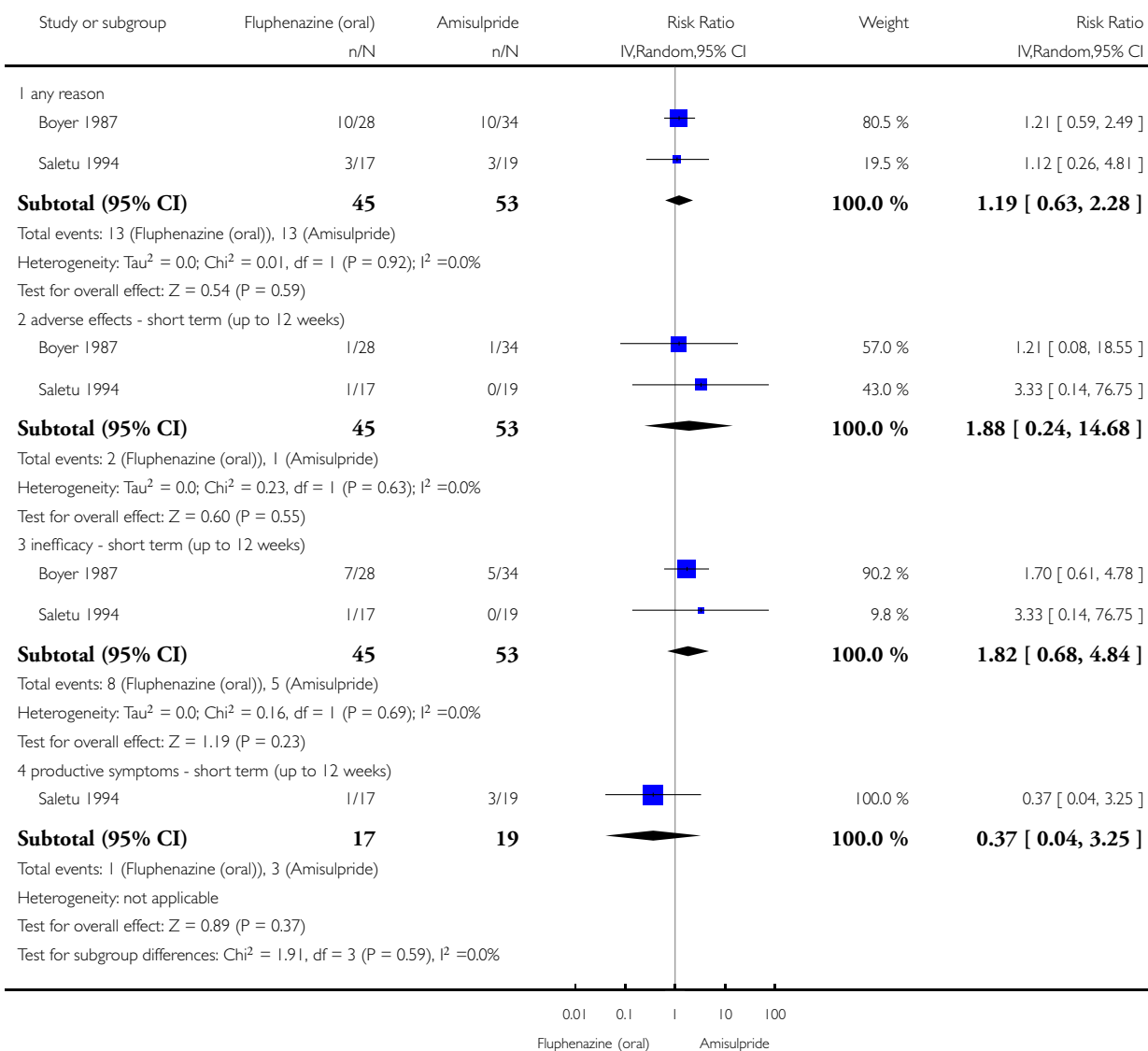


Analysis 1.5. Comparison 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE, Outcome 5 Leaving the study early.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 1 FLUPHENAZINE (ORAL) vs AMISULPRIDE

Outcome: 5 Leaving the study early

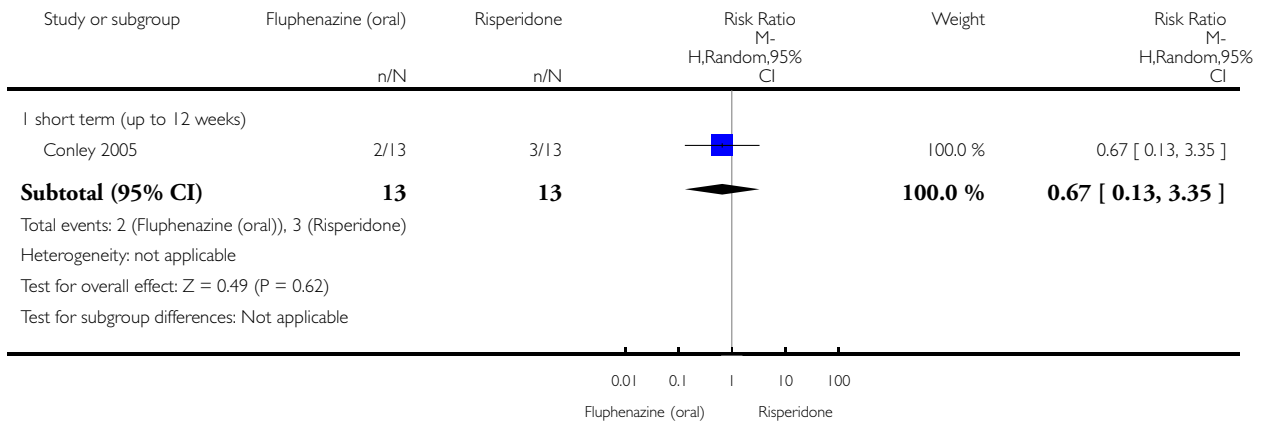


Analysis 2.1. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 1 Clinically important response (defined by study).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 1 Clinically important response (defined by study)

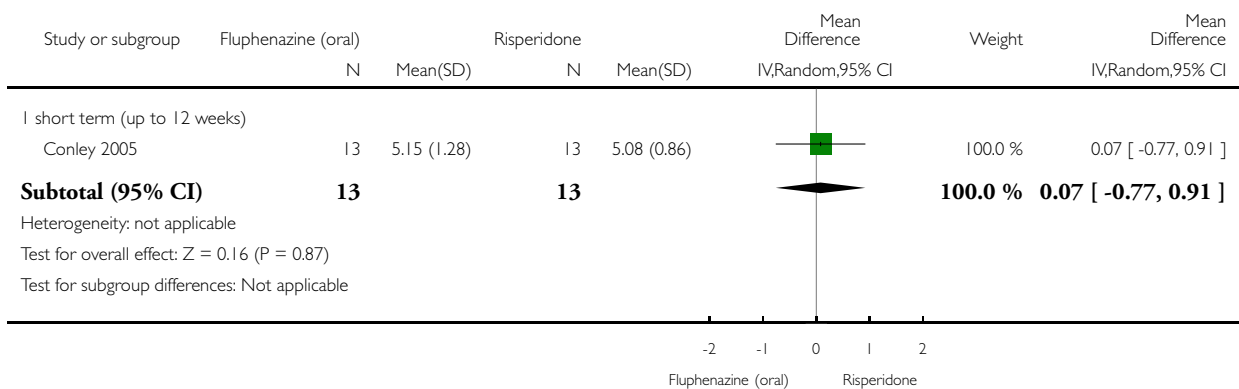


Analysis 2.2. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 2 Global state: I. Average endpoint score of CGI scales (high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 2 Global state: I. Average endpoint score of CGI scales (high = poor)

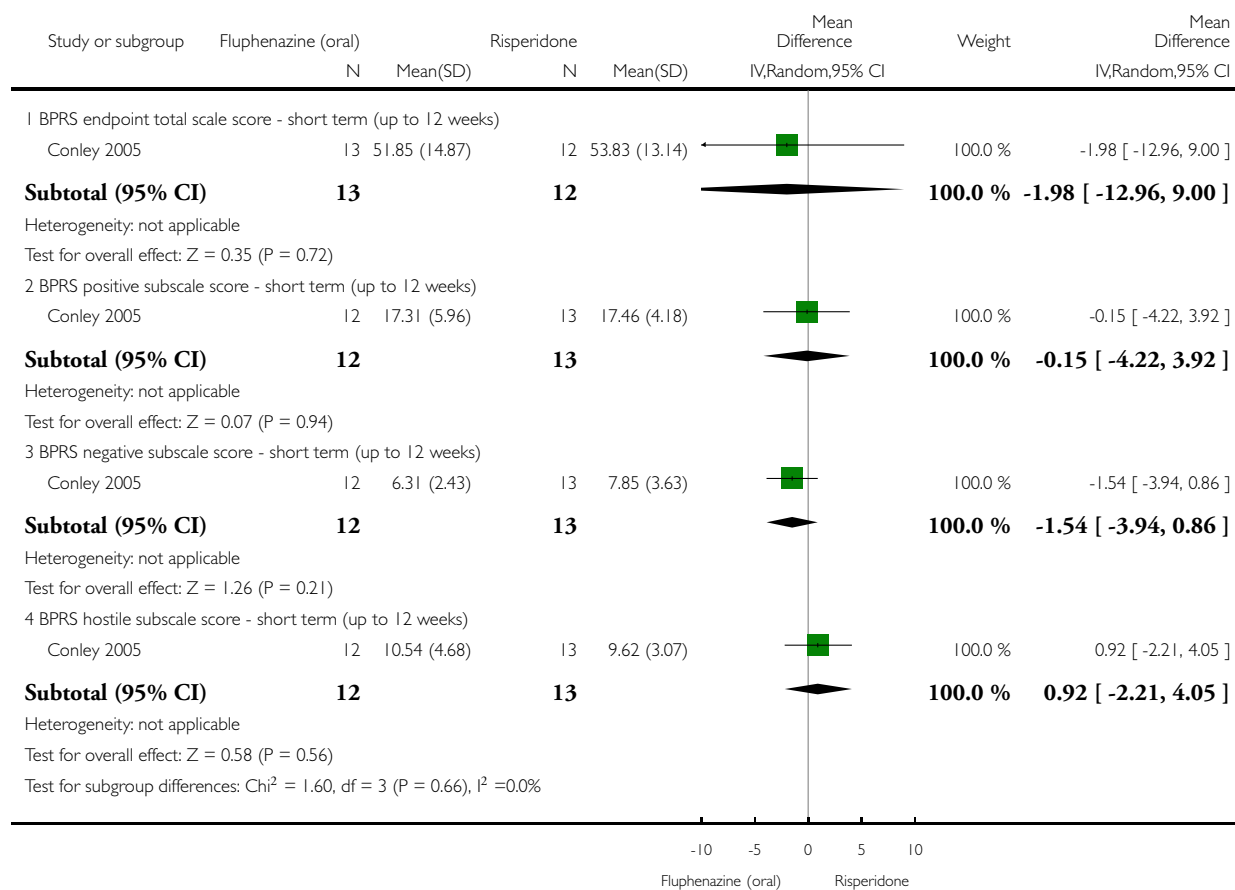


Analysis 2.3. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 3 Mental state: 2a. Average endpoint scores (BPRS, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 3 Mental state: 2a. Average endpoint scores (BPRS, high = poor)



Analysis 2.4. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 4 Mental state: 2b. Average endpoint score of various scales (high = poor) - short term (up to 12 weeks) (skewed data).

Mental state: 2b. Average endpoint score of various scales (high = poor) - short term (up to 12 weeks) (skewed data)

Study	Interventions	Mean	SD	N	BPRS anxiety/d
BPRS anxiety/depression subscale score					
Conley 2005	Fluphenazine	10.38	5.20	12	

Mental state: 2b. Average endpoint score of various scales (high = poor) - short term (up to 12 weeks) (skewed data) (Continued)

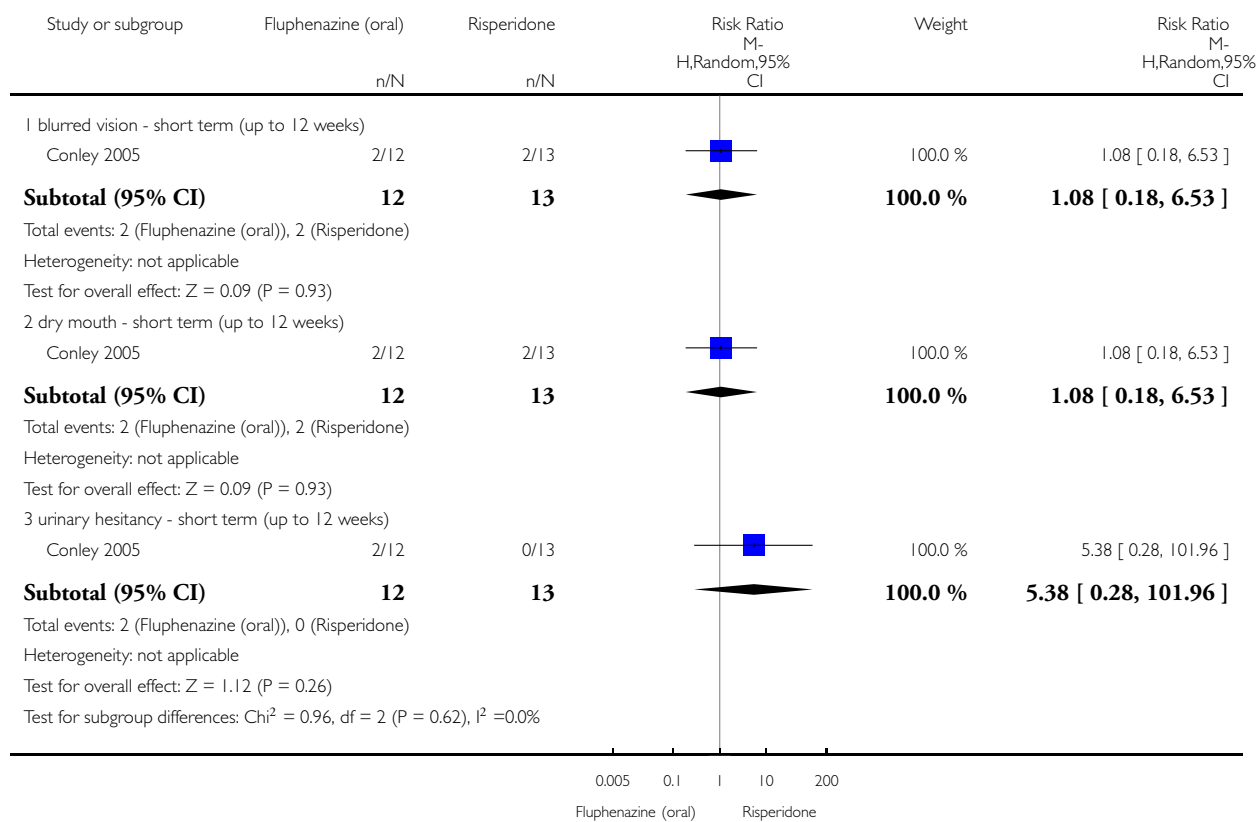
Conley 2005	Risperidone	9.31	5.91	13
BPRS activation subscale score				BPRS activation
Conley 2005	Fluphenazine	7.00	3.72	12
Conley 2005	Risperidone	7.00	3.06	13

Analysis 2.5. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 5 Adverse effects: 1. Anticholinergic effect.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 5 Adverse effects: 1. Anticholinergic effect

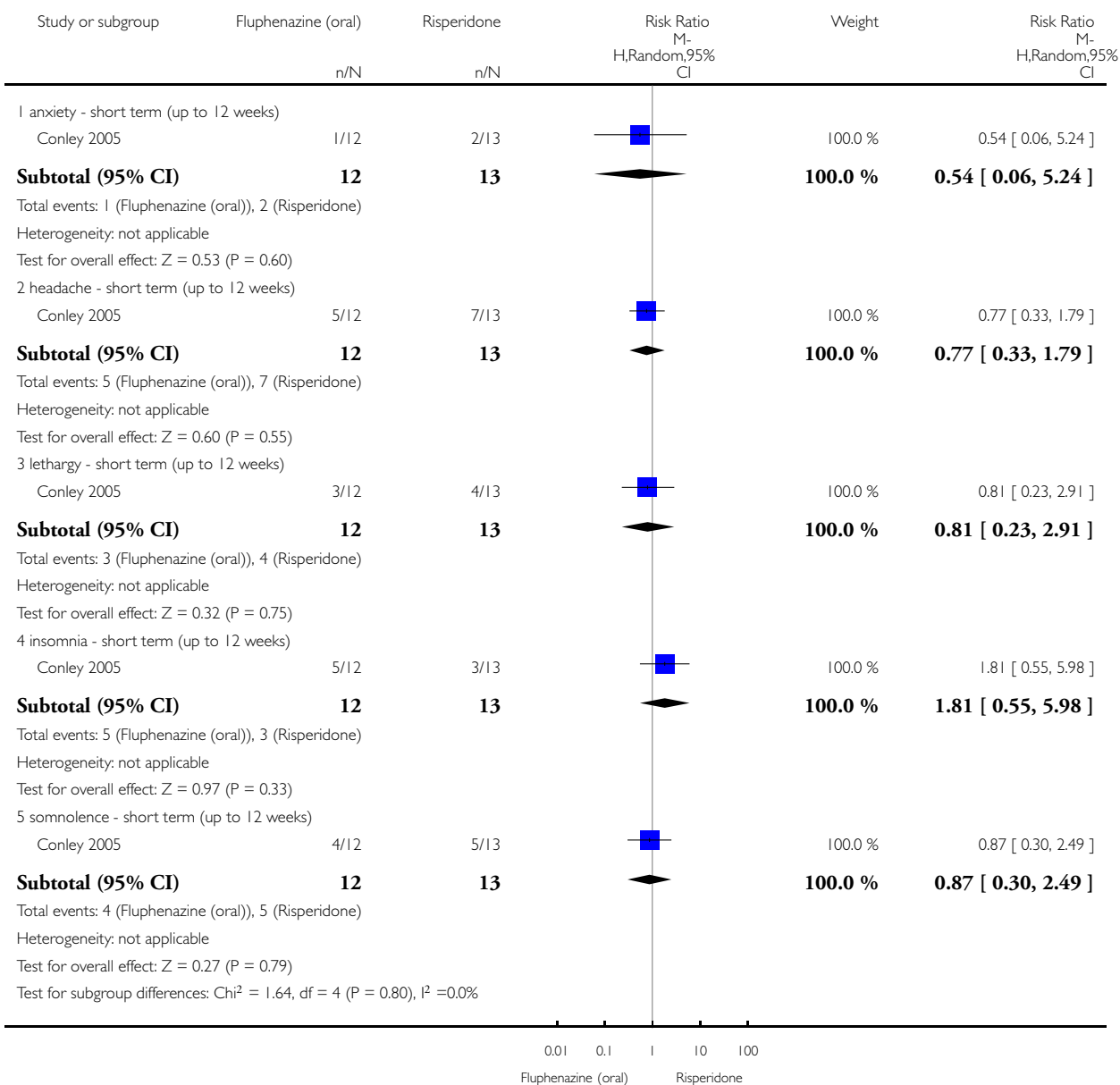


Analysis 2.6. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 6 Adverse effects: 2. Central nervous system.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 6 Adverse effects: 2. Central nervous system

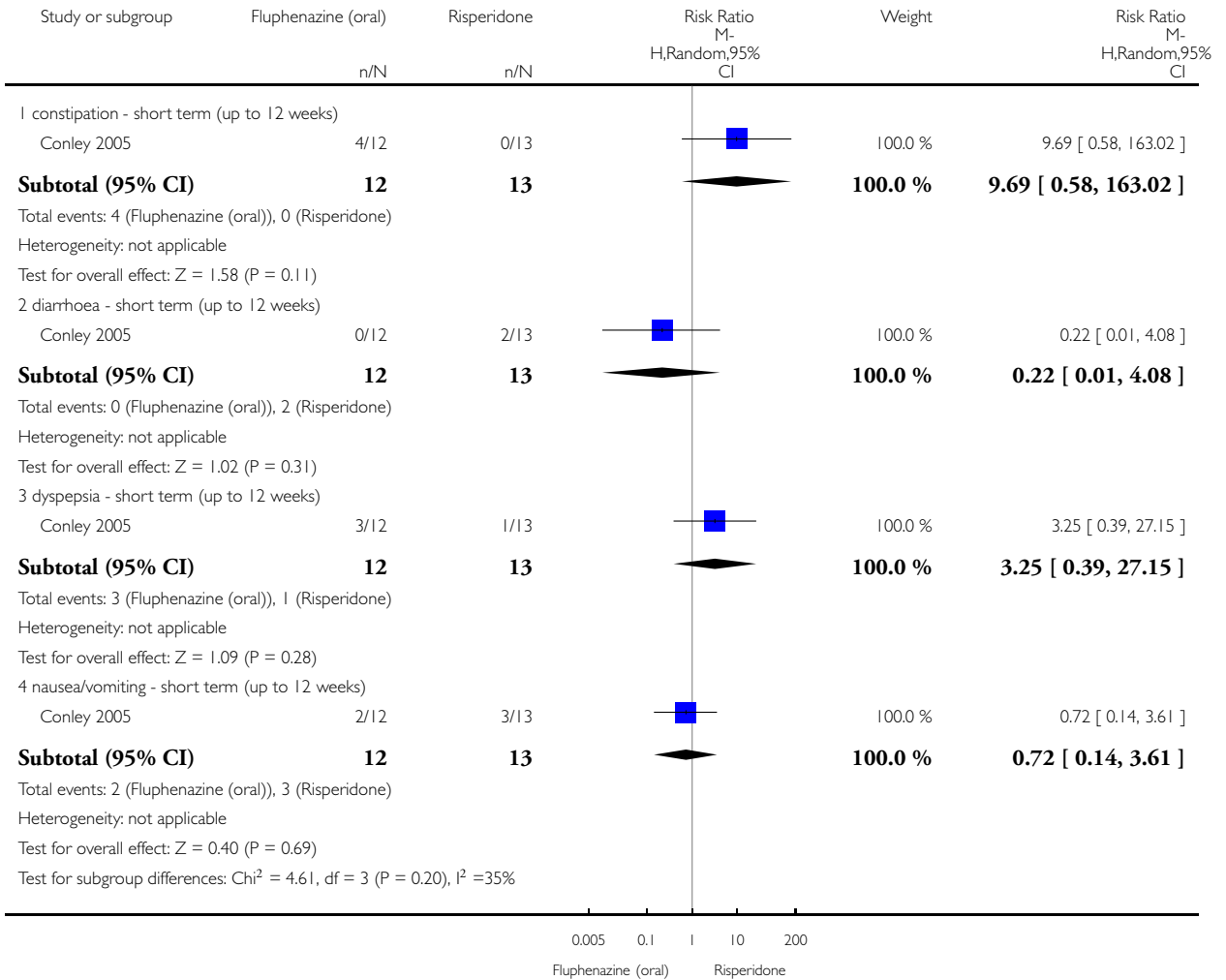


Analysis 2.7. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 7 Adverse effects: 3. Gastrointestinal.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 7 Adverse effects: 3. Gastrointestinal

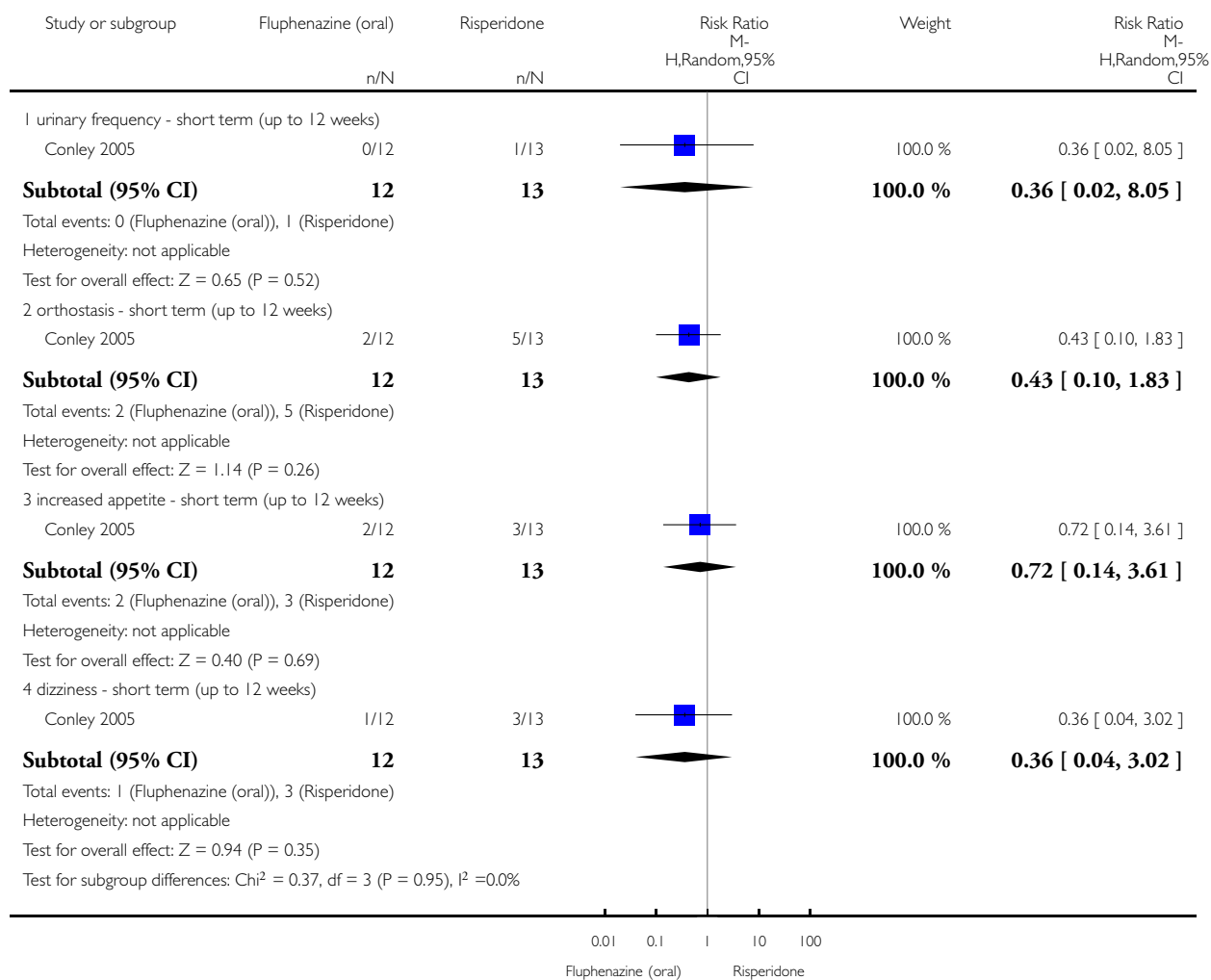


Analysis 2.8. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 8 Adverse effects: 4. Other adverse events.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 8 Adverse effects: 4. Other adverse events



Analysis 2.9. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 9 Adverse effects: 5. Average endpoint weight loss (kg) (skewed data).

Adverse effects: 5. Average endpoint weight loss (kg) (skewed data)

Study	Intervention	Mean(kg)	SD	N
short term (up to 12 weeks)				

Adverse effects: 5. Average endpoint weight loss (kg) (skewed data) (Continued)

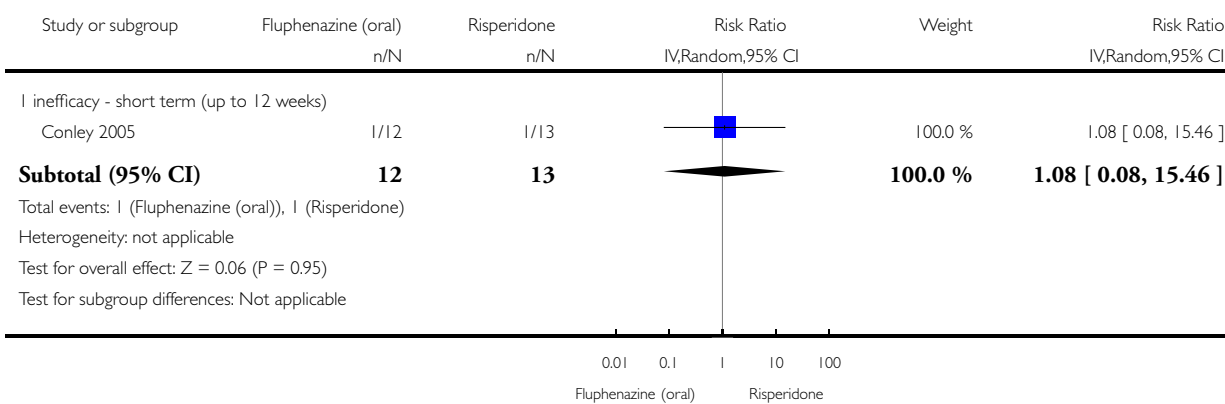
Conley 2005	Fluphenazine	-2.6	5.7	13
Conley 2005	Risperidone	-0.65	2.43	13

Analysis 2.10. Comparison 2 FLUPHENAZINE (ORAL) vs RISPERIDONE, Outcome 10 Leaving the study early.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 2 FLUPHENAZINE (ORAL) vs RISPERIDONE

Outcome: 10 Leaving the study early

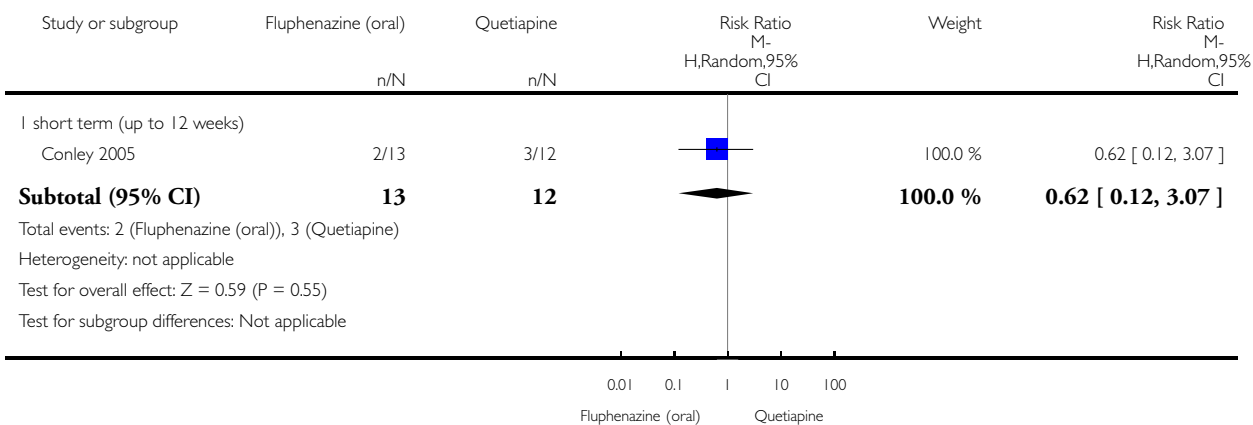


Analysis 3.1. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 1 Clinically important response (defined by study).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 1 Clinically important response (defined by study)

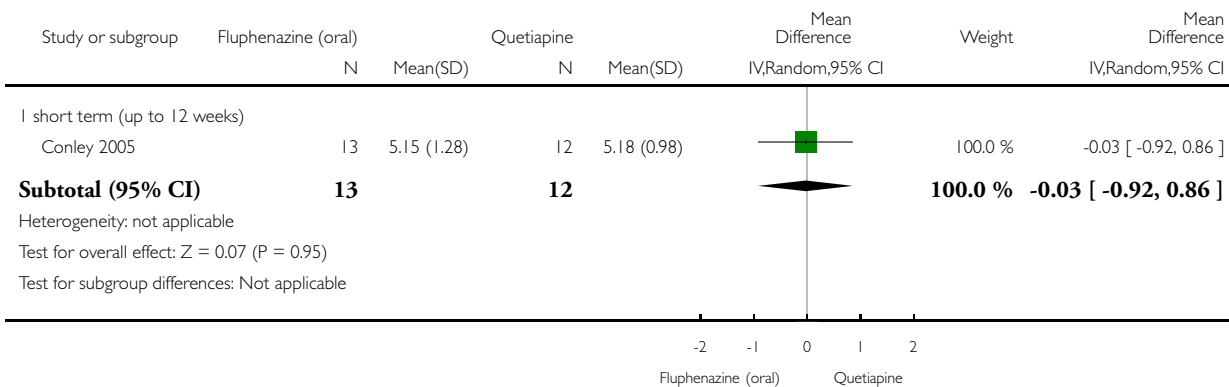


Analysis 3.2. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 2 Global state: 1. CGI: Average endpoint CGI-SI (high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

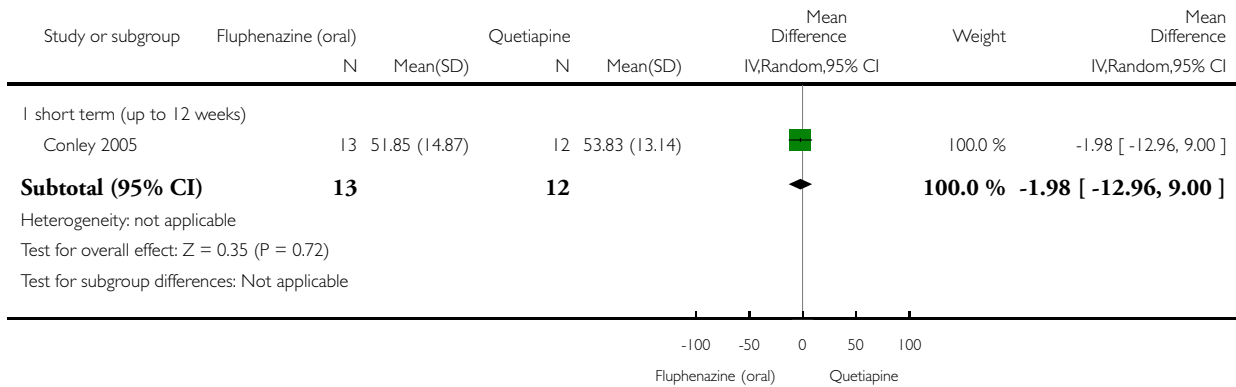
Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 2 Global state: 1. CGI: Average endpoint CGI-SI (high = poor)



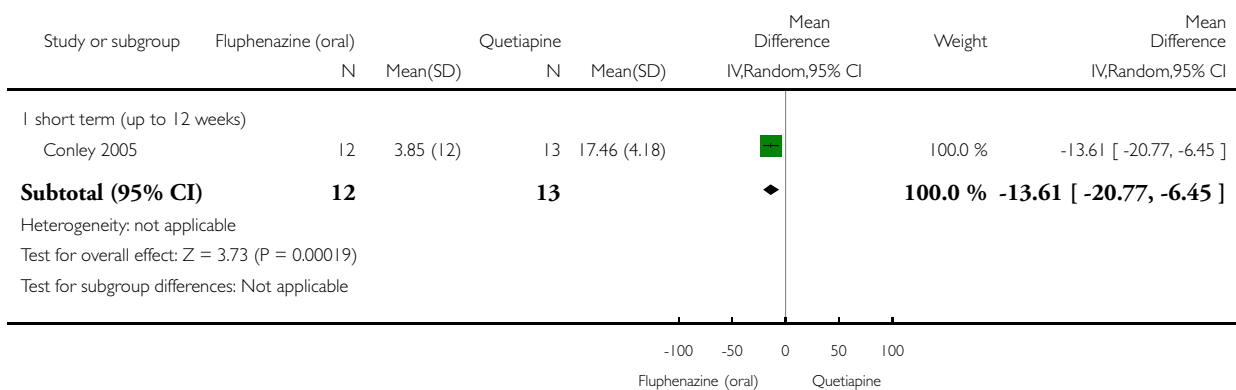
Analysis 3.3. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 3 Mental state: 2a. General - average endpoint score (BPRS total, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia
 Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE
 Outcome: 3 Mental state: 2a. General - average endpoint score (BPRS total, high = poor)



Analysis 3.4. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 4 Mental state: 2b. Positive symptoms - average endpoint score (BPRS positive sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia
 Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE
 Outcome: 4 Mental state: 2b. Positive symptoms - average endpoint score (BPRS positive sub-score, high = poor)

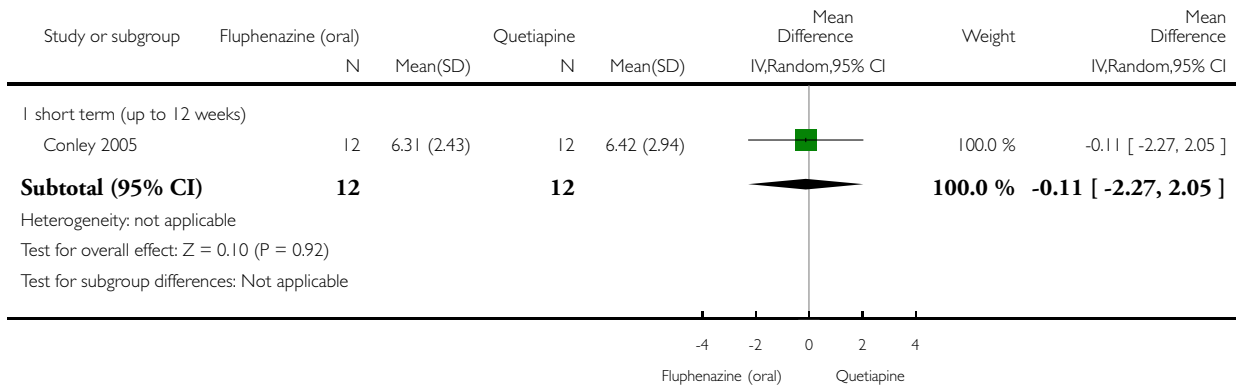


Analysis 3.5. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 5 Mental state: 2c. Negative symptoms - average endpoint score (BPRS negative sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 5 Mental state: 2c. Negative symptoms - average endpoint score (BPRS negative sub-score, high = poor)



Analysis 3.6. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 6 Mental state: 2d. Anxiety/depression symptoms - average endpoint score (BPRS anxiety/depression sub-score, high score = poor).

Mental state: 2d. Anxiety/depression symptoms - average endpoint score (BPRS anxiety/depression sub-score, high score = poor)

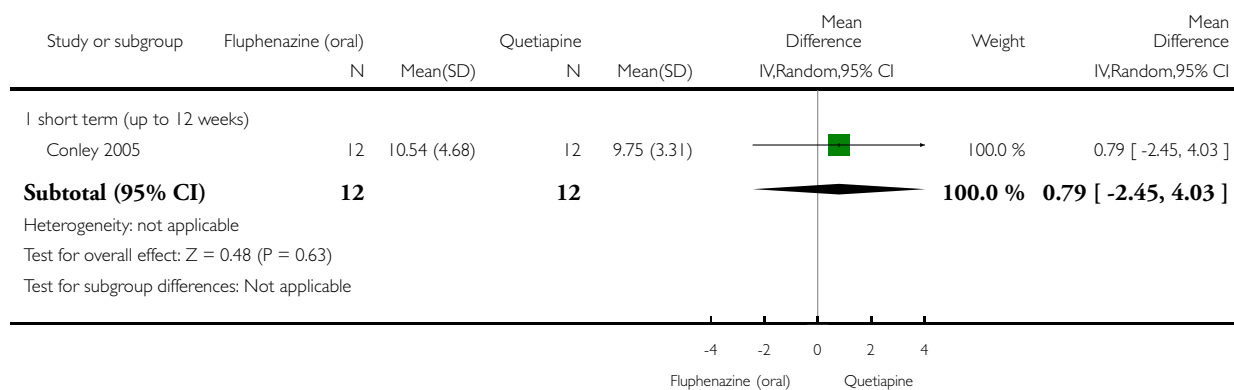
Study	Interventions	Mean	SD	N
short term (up to 12 weeks)				
Conley 2005	Quetiapine	10.25	3.70	12
Conley 2005	Risperidone	9.31	5.91	13

Analysis 3.7. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 7 Mental state: 2e. Hostility symptoms - average endpoint score (BPRS hostility sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 7 Mental state: 2e. Hostility symptoms - average endpoint score (BPRS hostility sub-score, high = poor)



Analysis 3.8. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 8 Mental state: 2f. Activation symptoms - average endpoint score (BPRS activation sub-score, high = poor).

Mental state: 2f. Activation symptoms - average endpoint score (BPRS activation sub-score, high = poor)

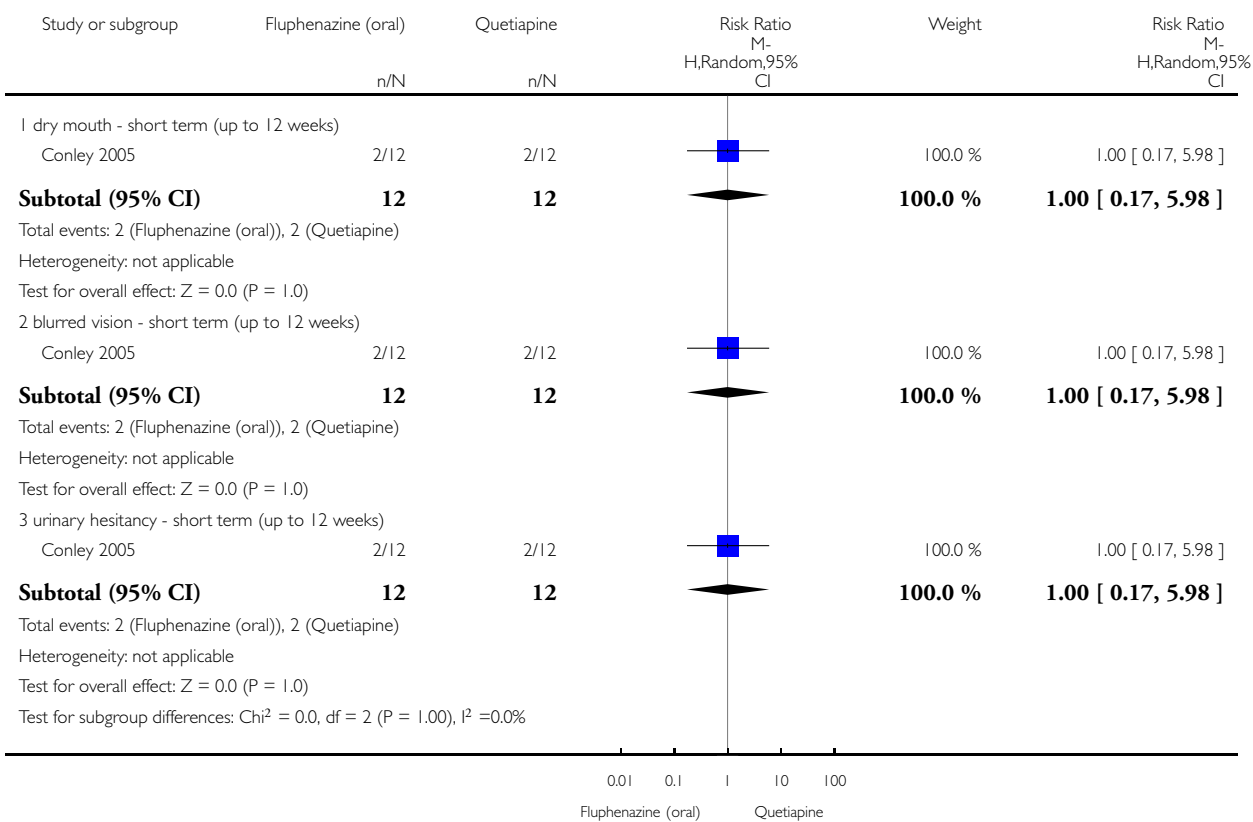
Study	Intervention	Mean	SD	N
short term (up to 12 weeks)				
Conley 2005	Quetiapine	7.33	2.77	12
Conley 2005	Risperidone	7.00	3.06	13

Analysis 3.9. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 9 Adverse effects: 1. Anticholinergic effect.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 9 Adverse effects: 1. Anticholinergic effect

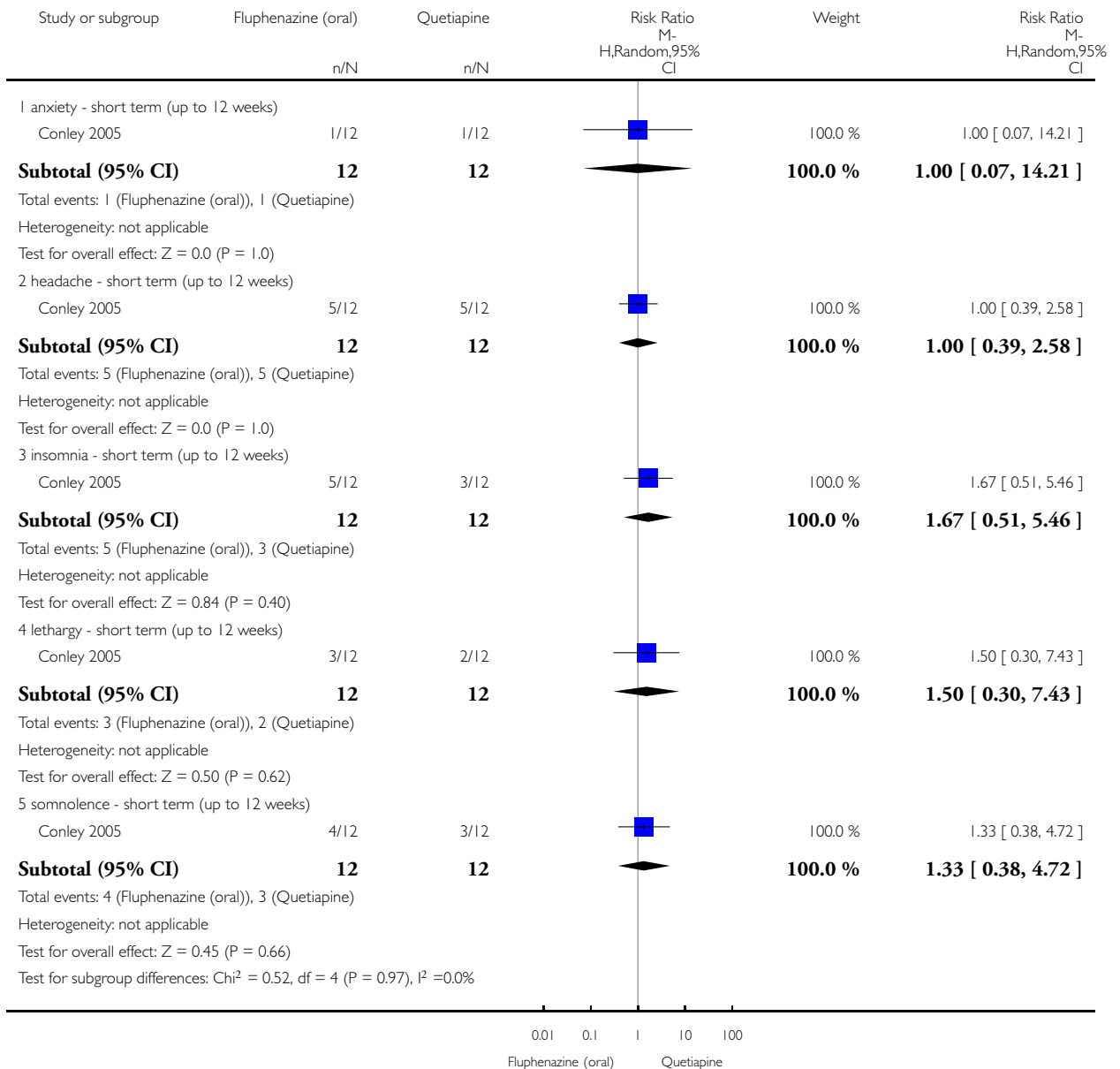


Analysis 3.10. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 10 Adverse effects: 2. Central nervous system.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 10 Adverse effects: 2. Central nervous system

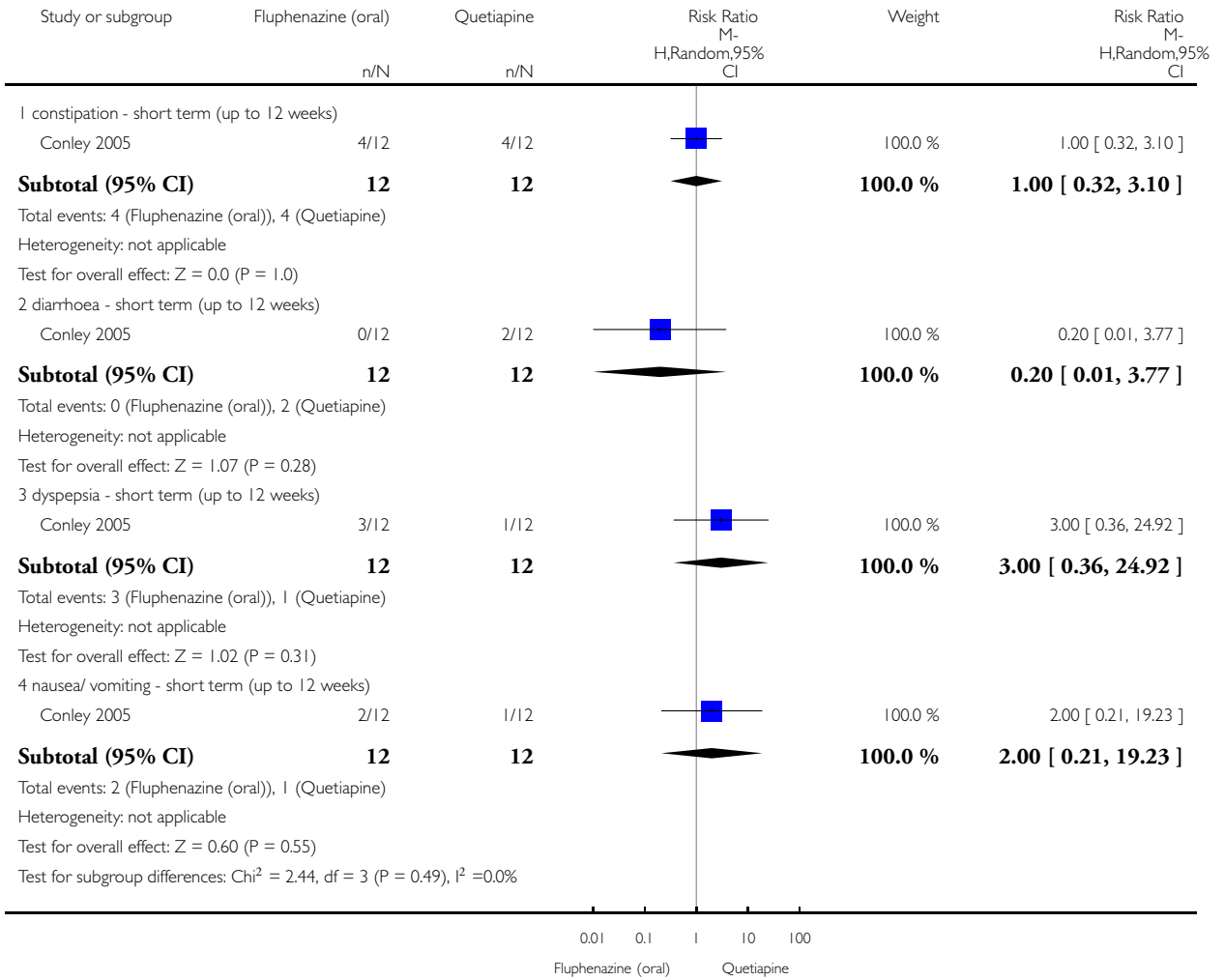


Analysis 3.11. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 1 | Adverse effects: 3. Gastrointestinal adverse effects.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 1 | Adverse effects: 3. Gastrointestinal adverse effects

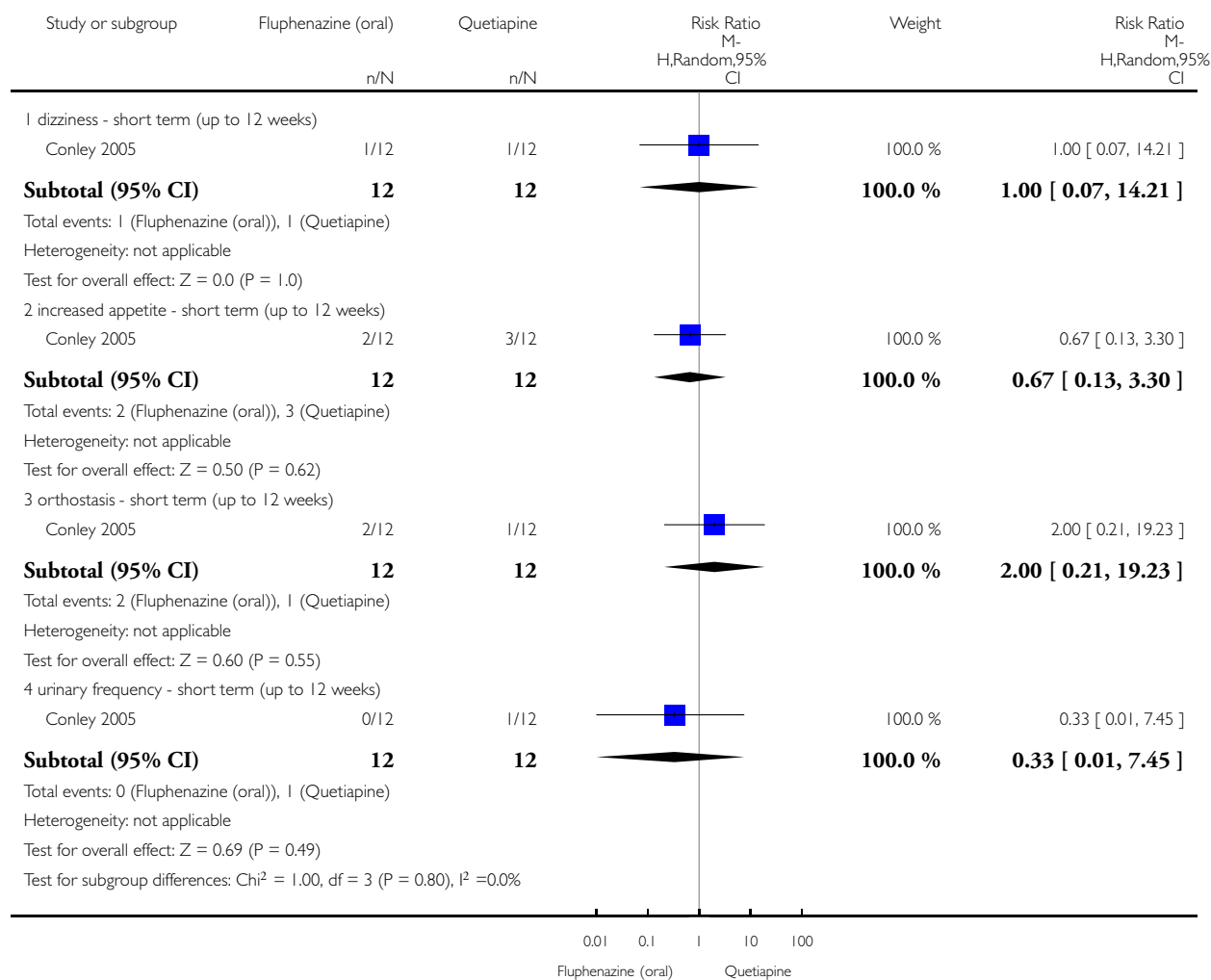


Analysis 3.12. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 12 Adverse effects: 4a. Other adverse events.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 12 Adverse effects: 4a. Other adverse events



Analysis 3.13. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 13 Adverse effects: 4b. Other - average endpoint weight loss (average weight in kg) (skewed).

Adverse effects: 4b. Other - average endpoint weight loss (average weight in kg) (skewed)

Study	Intervention	Mean(kg)	SD	N
short term (up to 12 weeks)				

Fluphenazine (oral) versus atypical antipsychotics for schizophrenia (Review)

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Adverse effects: 4b. Other - average endpoint weight loss (average weight in kg) (skewed) (Continued)

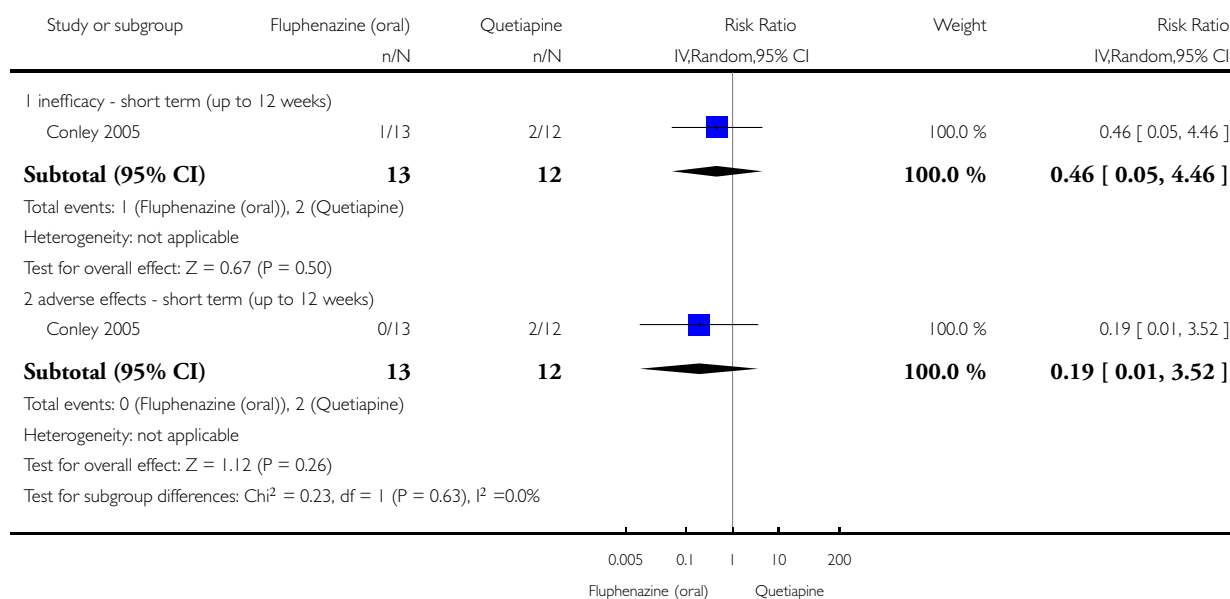
Conley 2005	Fluphenazine	-2.6	5.7	13
Conley 2005	Risperidone	-0.65	2.43	13

Analysis 3.14. Comparison 3 FLUPHENAZINE (ORAL) vs QUETIAPINE, Outcome 14 Leaving the study early.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 3 FLUPHENAZINE (ORAL) vs QUETIAPINE

Outcome: 14 Leaving the study early

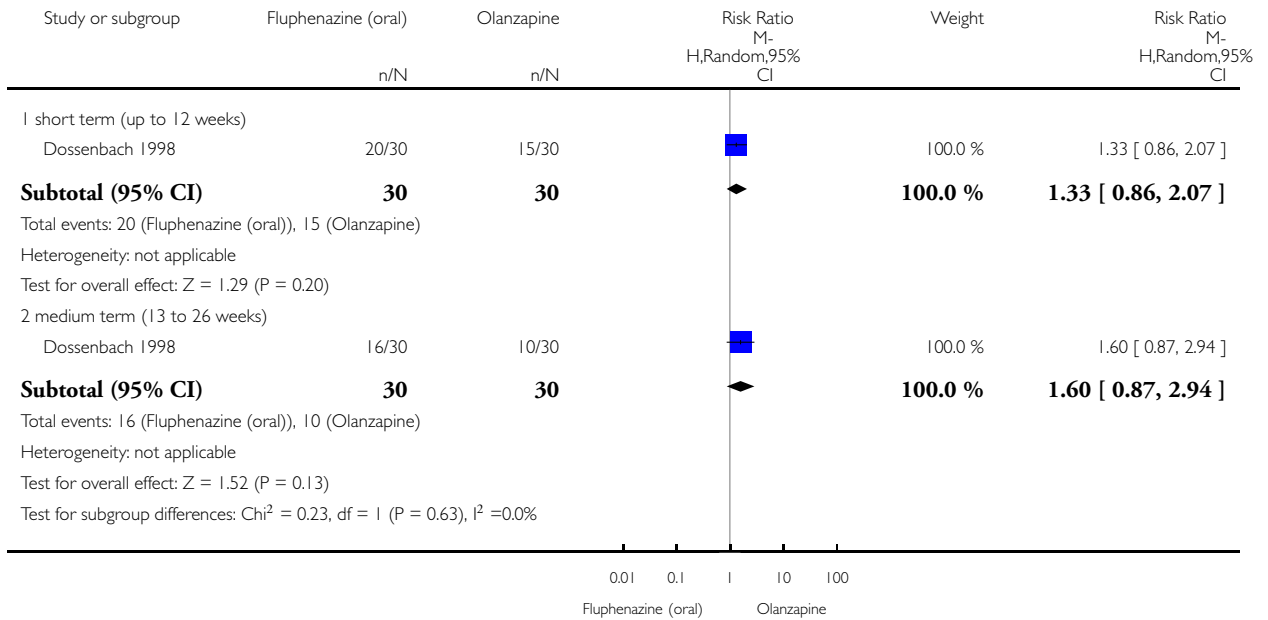


Analysis 4.1. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 1 Clinically important response (defined by author).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 1 Clinically important response (defined by author)

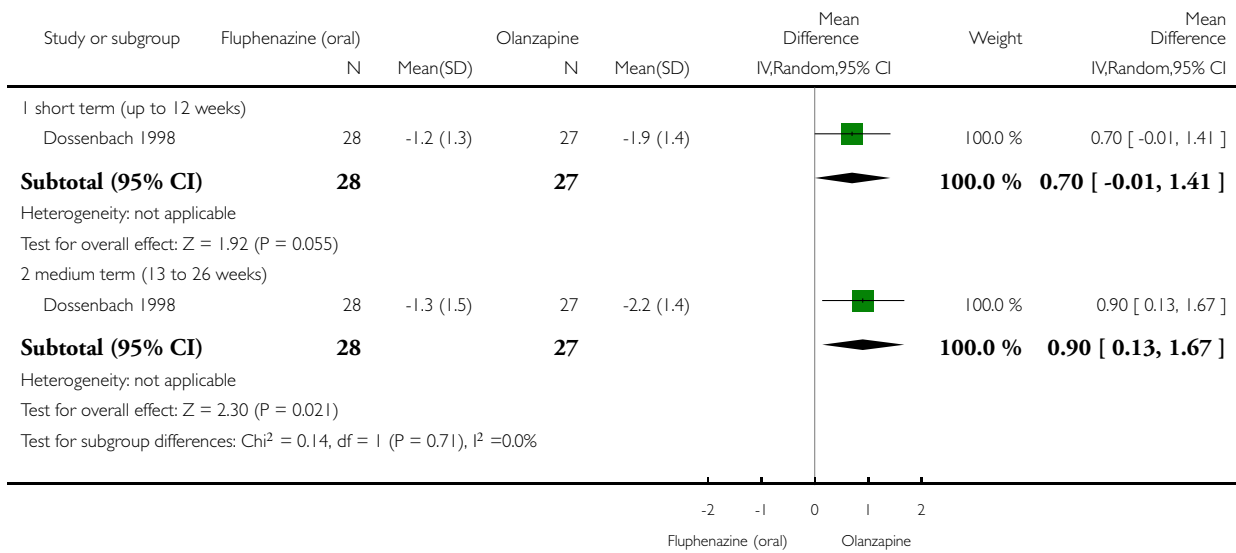


Analysis 4.2. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 2 Global state: I. CGI: Average change CGI-SI (high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 2 Global state: I. CGI: Average change CGI-SI (high = poor)

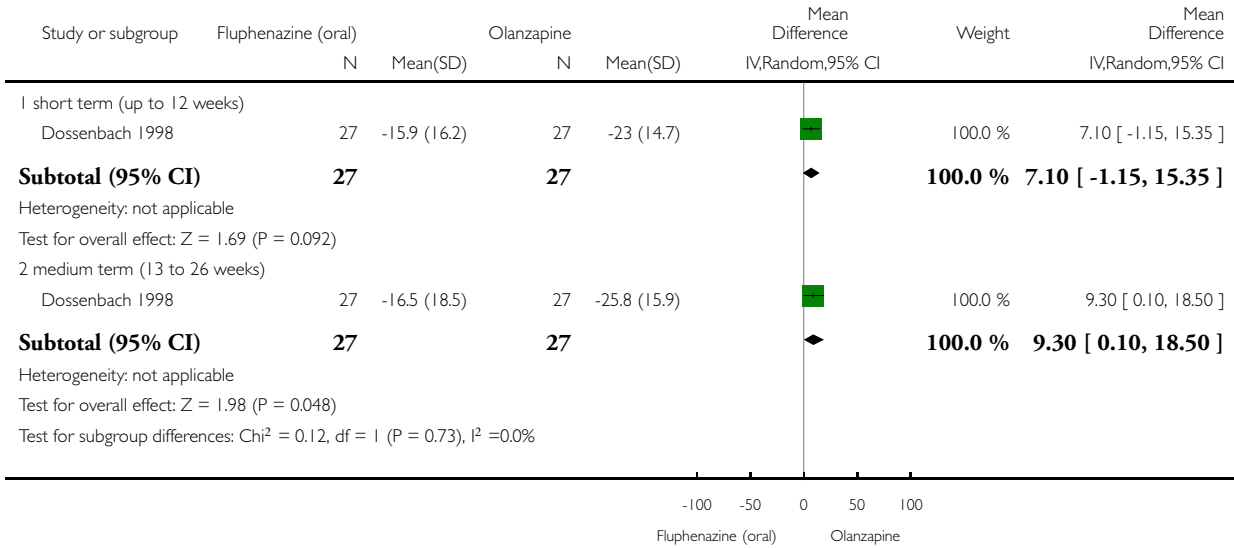


Analysis 4.3. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 3 Mental state: 2a. General - average change score (BPRS total, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 3 Mental state: 2a. General - average change score (BPRS total, high = poor)

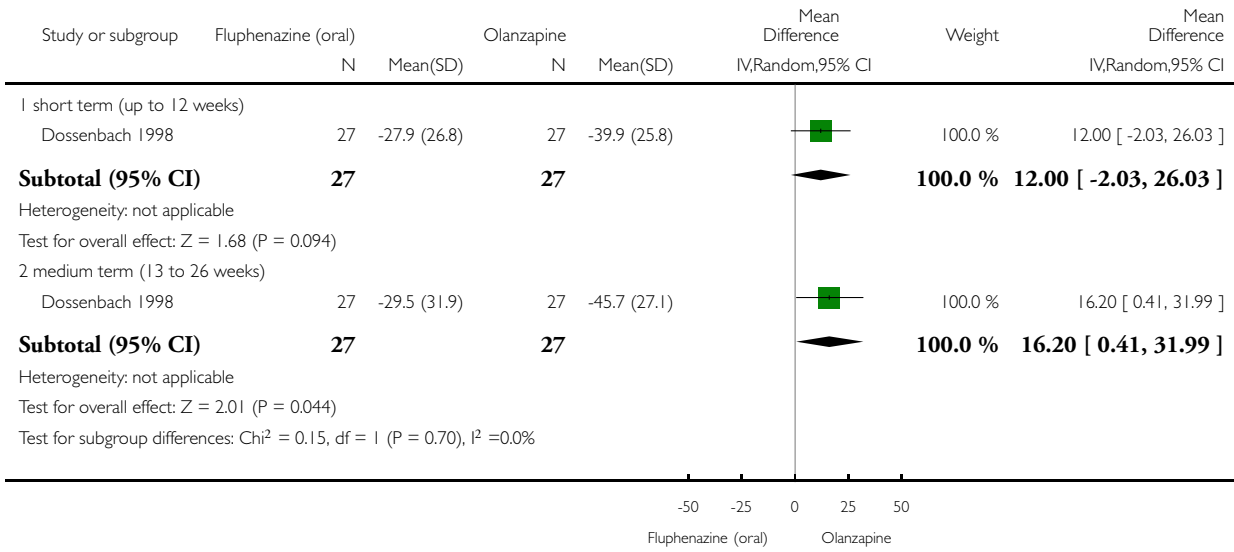


Analysis 4.4. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 4 Mental state: 2b. General - average change score (PANSS total, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 4 Mental state: 2b. General - average change score (PANSS total, high = poor)

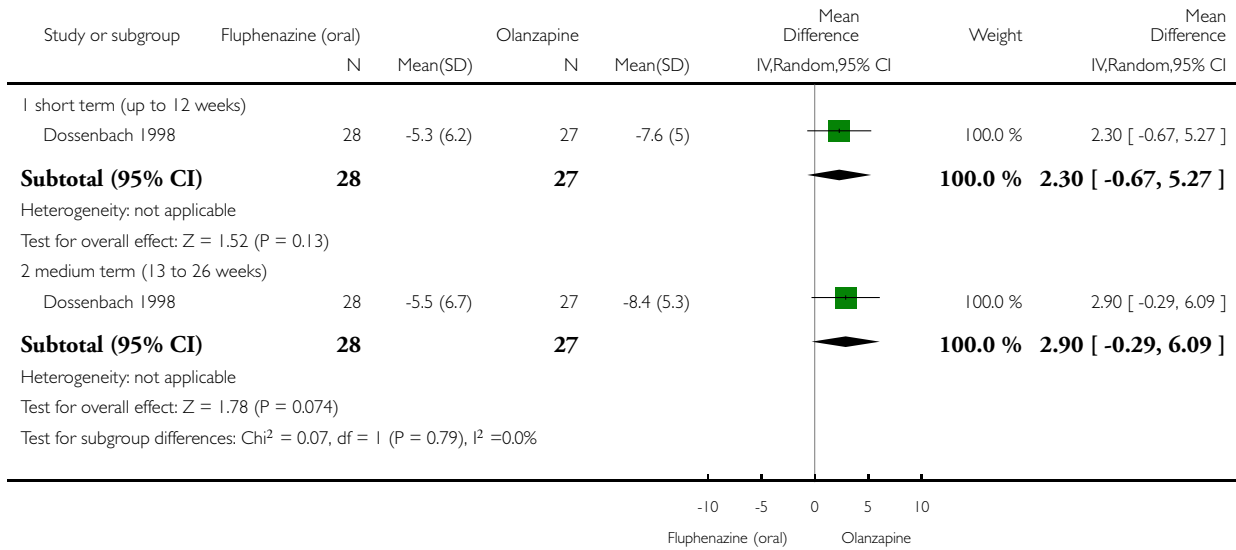


Analysis 4.5. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 5 Mental state: 2c. Positive symptoms - average change score (BPRS positive sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 5 Mental state: 2c. Positive symptoms - average change score (BPRS positive sub-score, high = poor)

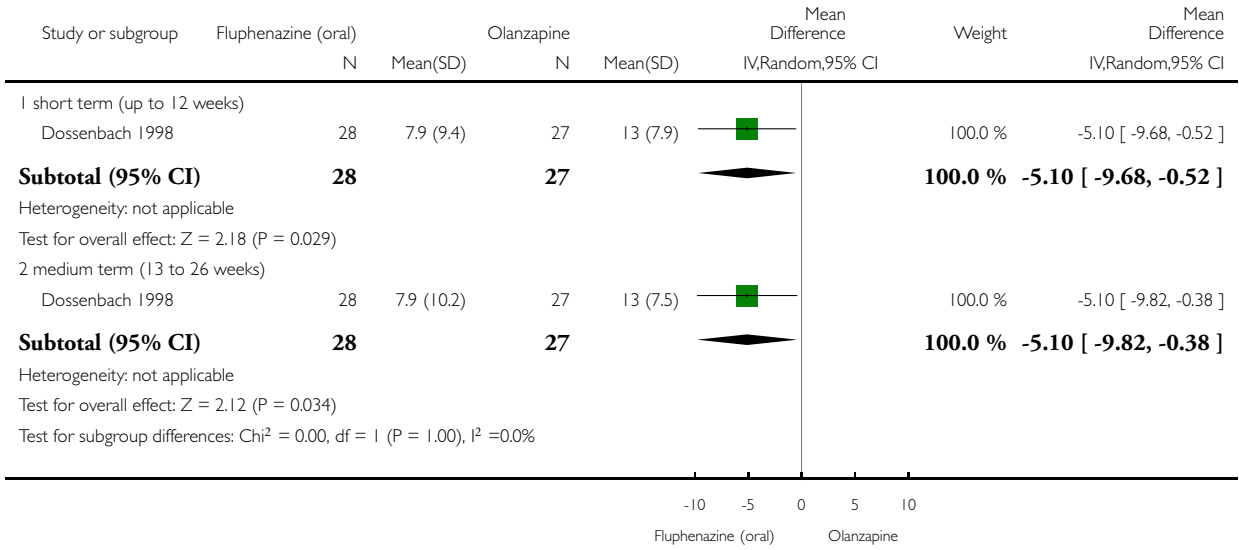


Analysis 4.6. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 6 Mental state: 2d. Positive symptoms - average endpoint score (PANSS positive sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 6 Mental state: 2d. Positive symptoms - average endpoint score (PANSS positive sub-score, high = poor)

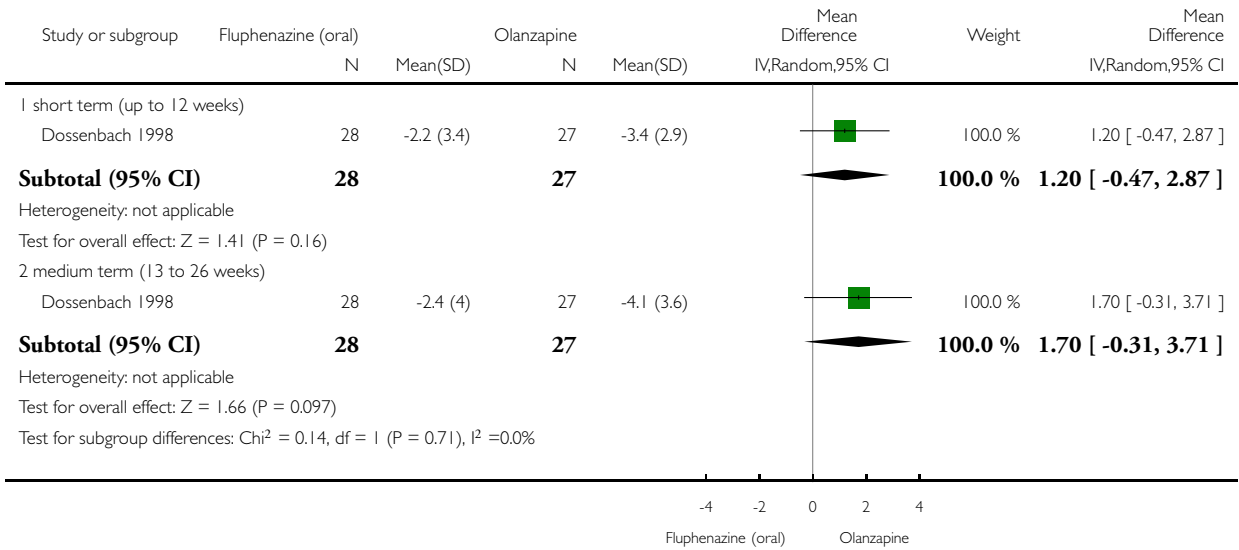


Analysis 4.7. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 7 Mental state: 2e. Negative symptoms - average change score (BPRS negative sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 7 Mental state: 2e. Negative symptoms - average change score (BPRS negative sub-score, high = poor)

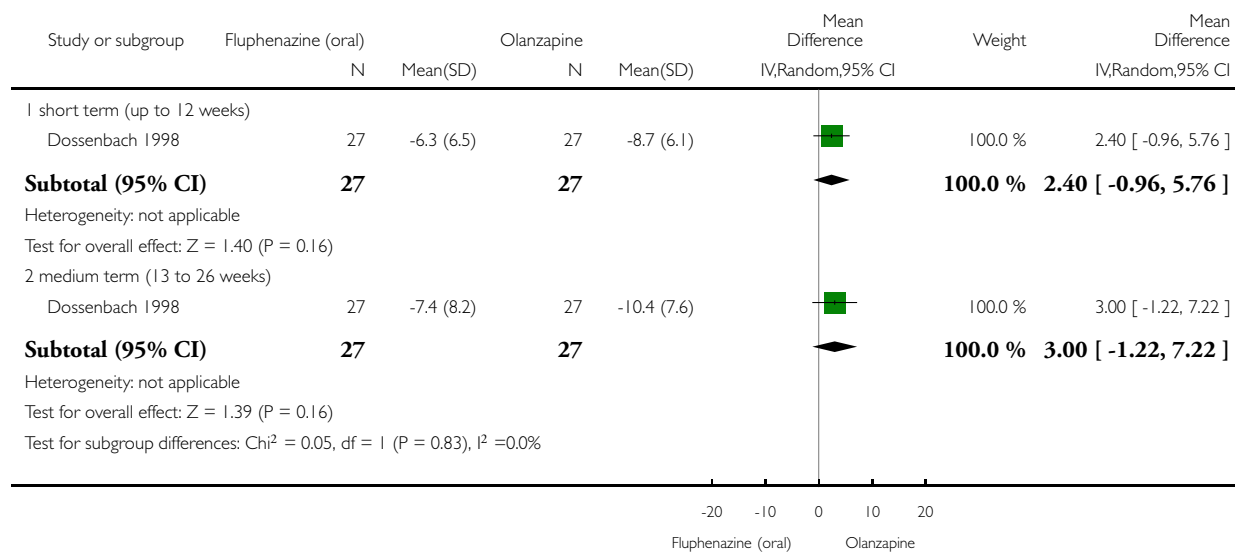


Analysis 4.8. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 8 Mental state: 2f. Negative symptoms - average change score (PANSS negative sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 8 Mental state: 2f. Negative symptoms - average change score (PANSS negative sub-score, high = poor)

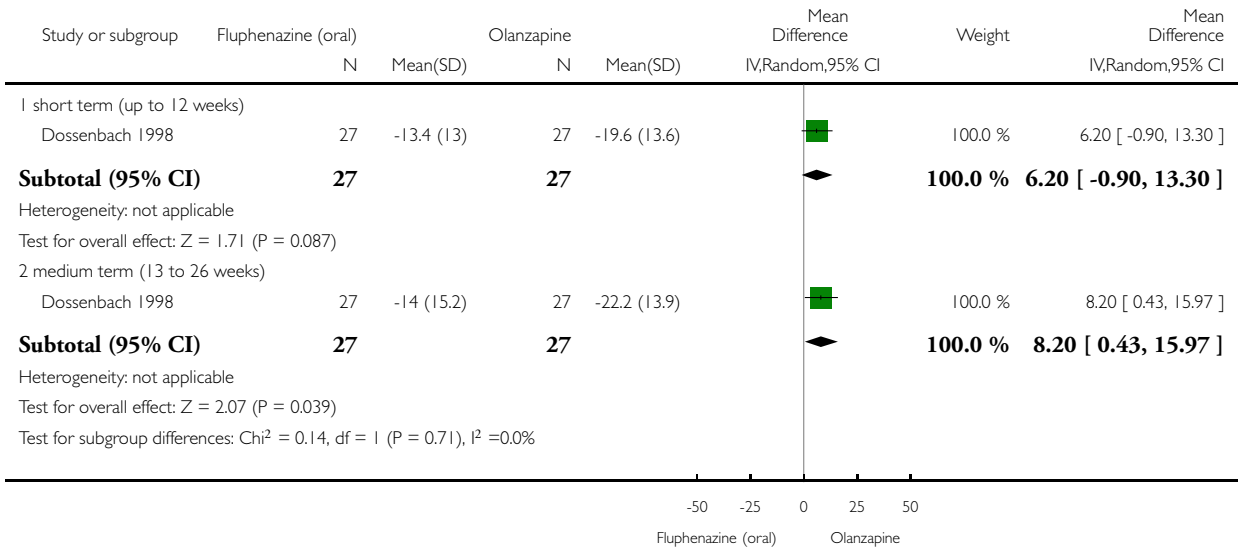


Analysis 4.9. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 9 Mental state: 2g. General psychopathology - average change score (PANSS general psychopathology sub-score, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 9 Mental state: 2g. General psychopathology - average change score (PANSS general psychopathology sub-score, high = poor)

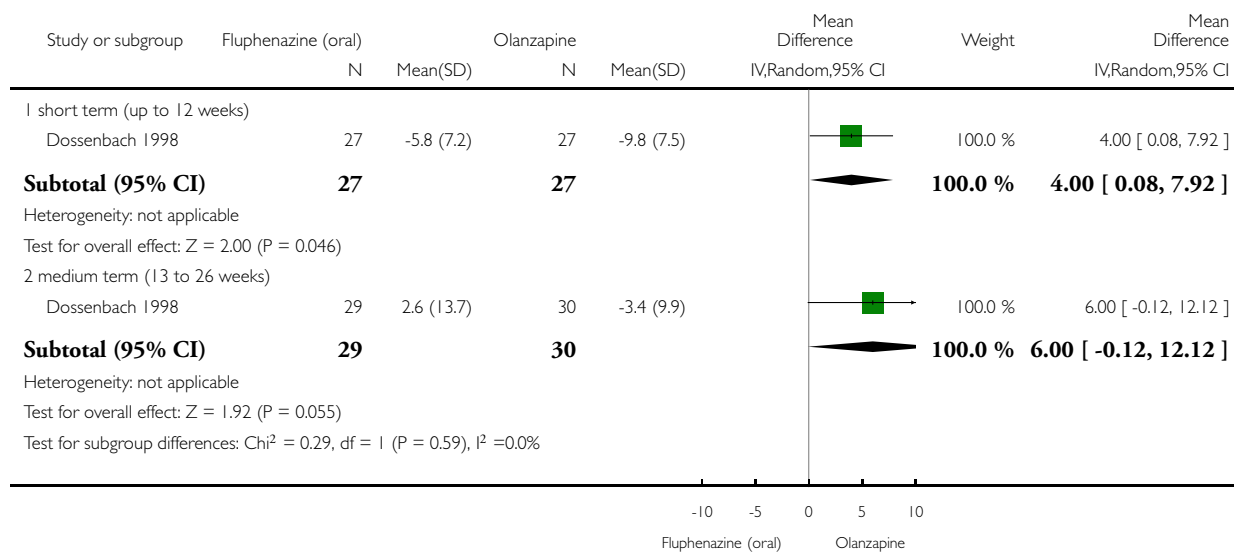


Analysis 4.10. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 10 Mental state: 2h. Anxiety - average change score (HAMA, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 10 Mental state: 2h. Anxiety - average change score (HAMA, high = poor)

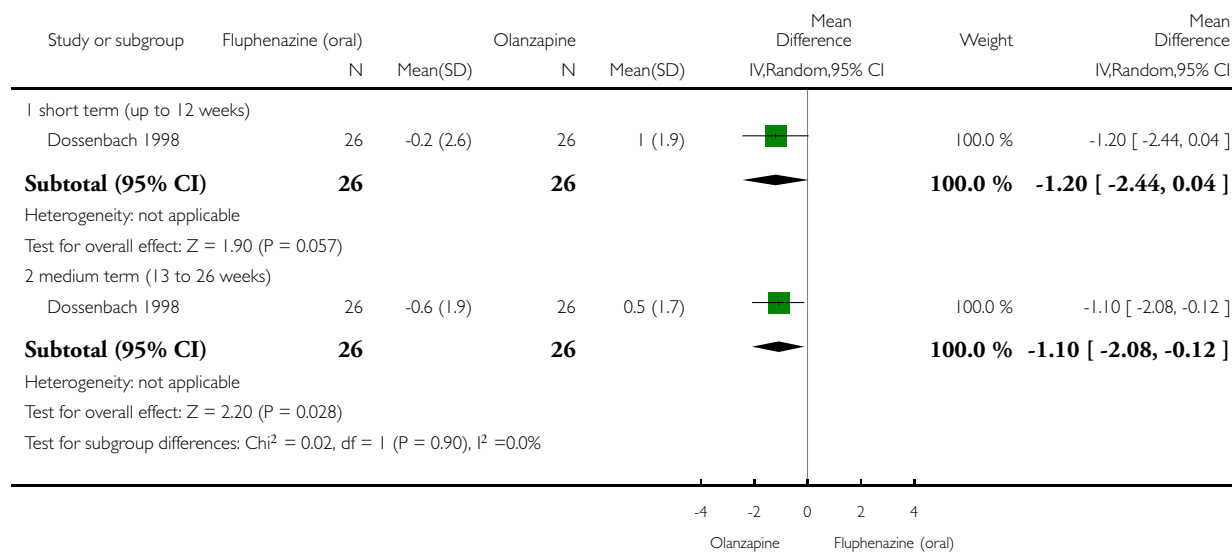


Analysis 4.11. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 11 Satisfaction with treatment: 1. Average change score (DAI, low = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 11 Satisfaction with treatment: 1. Average change score (DAI, low = poor)

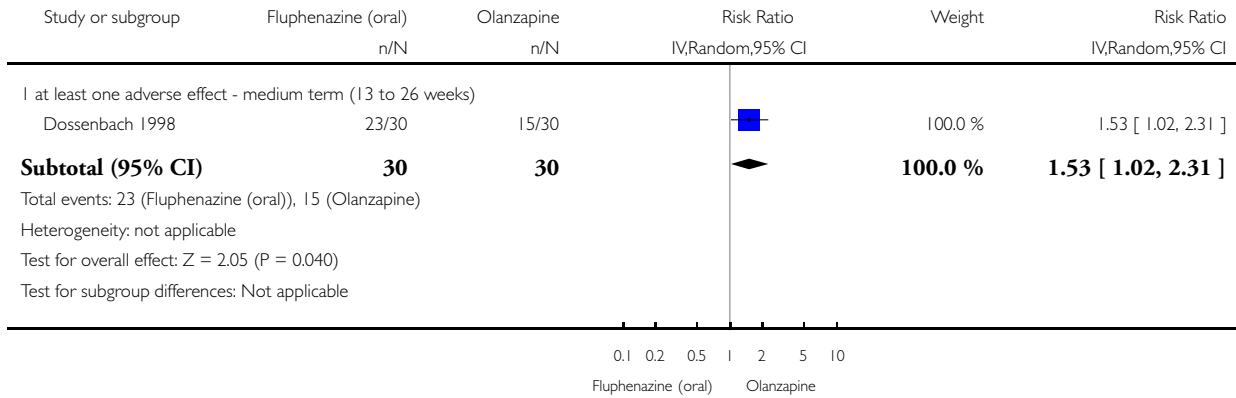


Analysis 4.12. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 12 Adverse effects: 1. General.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 12 Adverse effects: 1. General

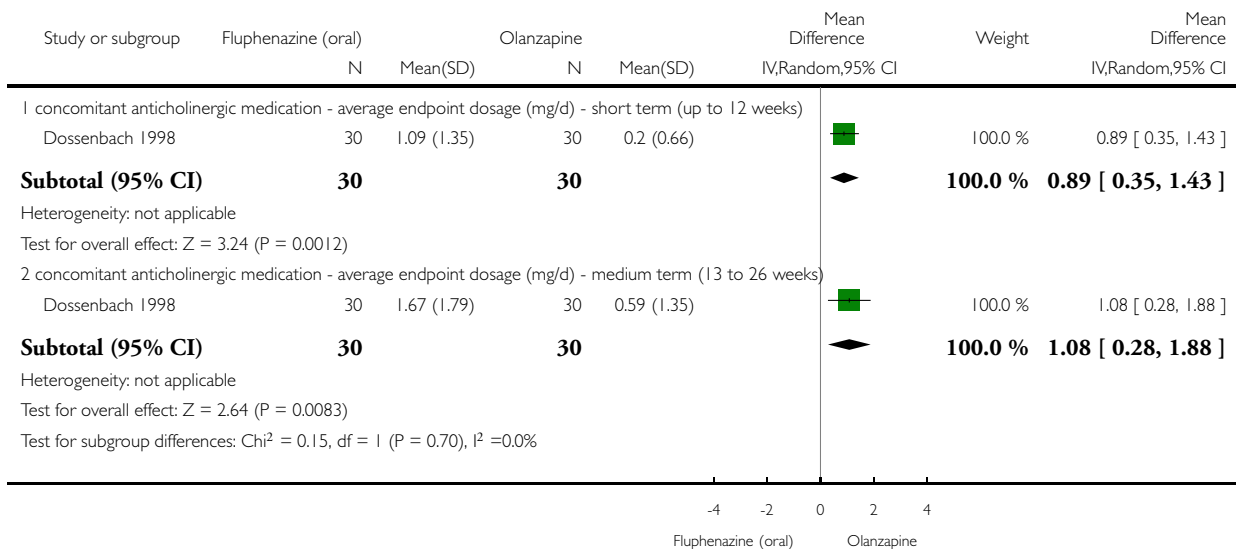


Analysis 4.13. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 13 Adverse effects: 2. Anticholinergic effect.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 13 Adverse effects: 2. Anticholinergic effect

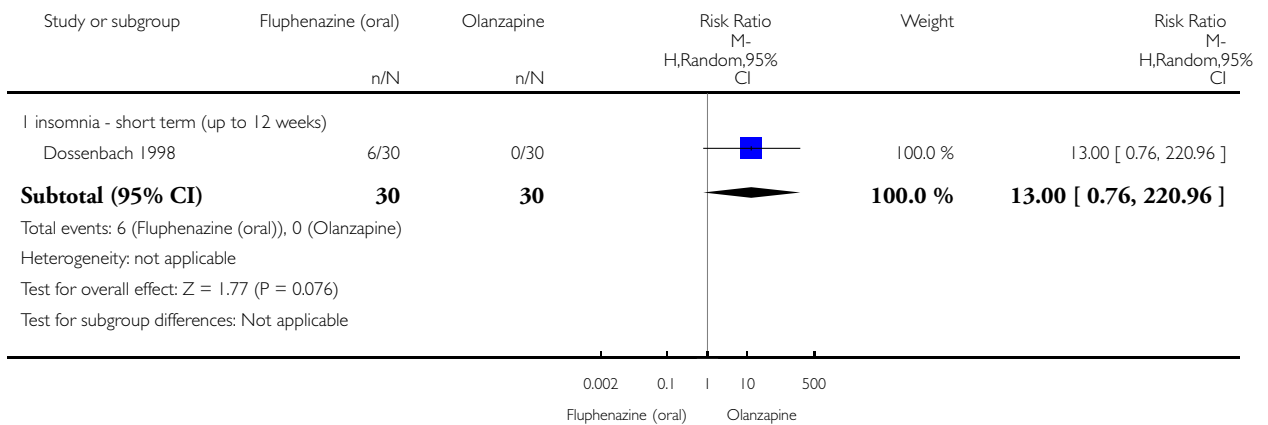


Analysis 4.14. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 14 Adverse effects: 3a. Central nervous system.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 14 Adverse effects: 3a. Central nervous system

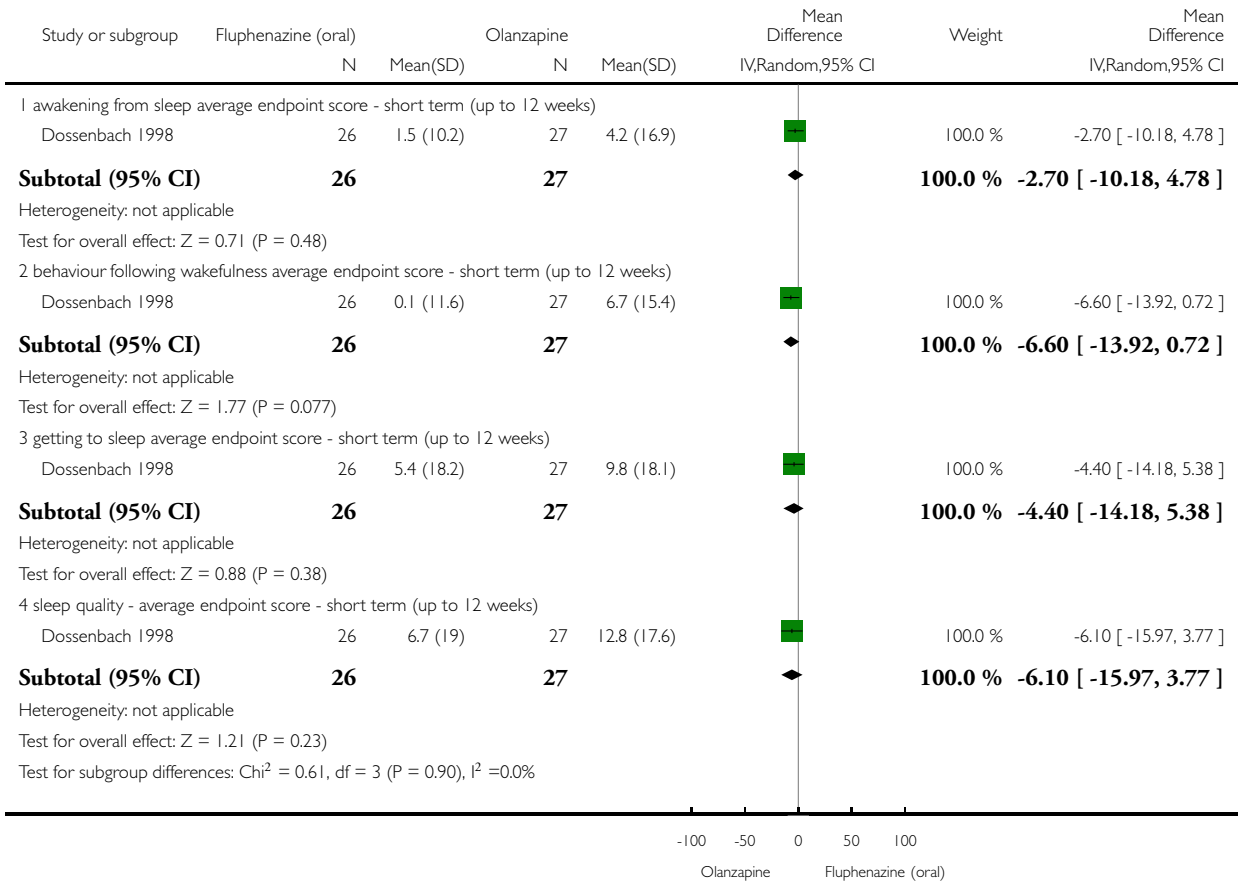


Analysis 4.15. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 15 Adverse effects: 3b. CNS (LSEQ, low = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 15 Adverse effects: 3b. CNS (LSEQ, low = poor)

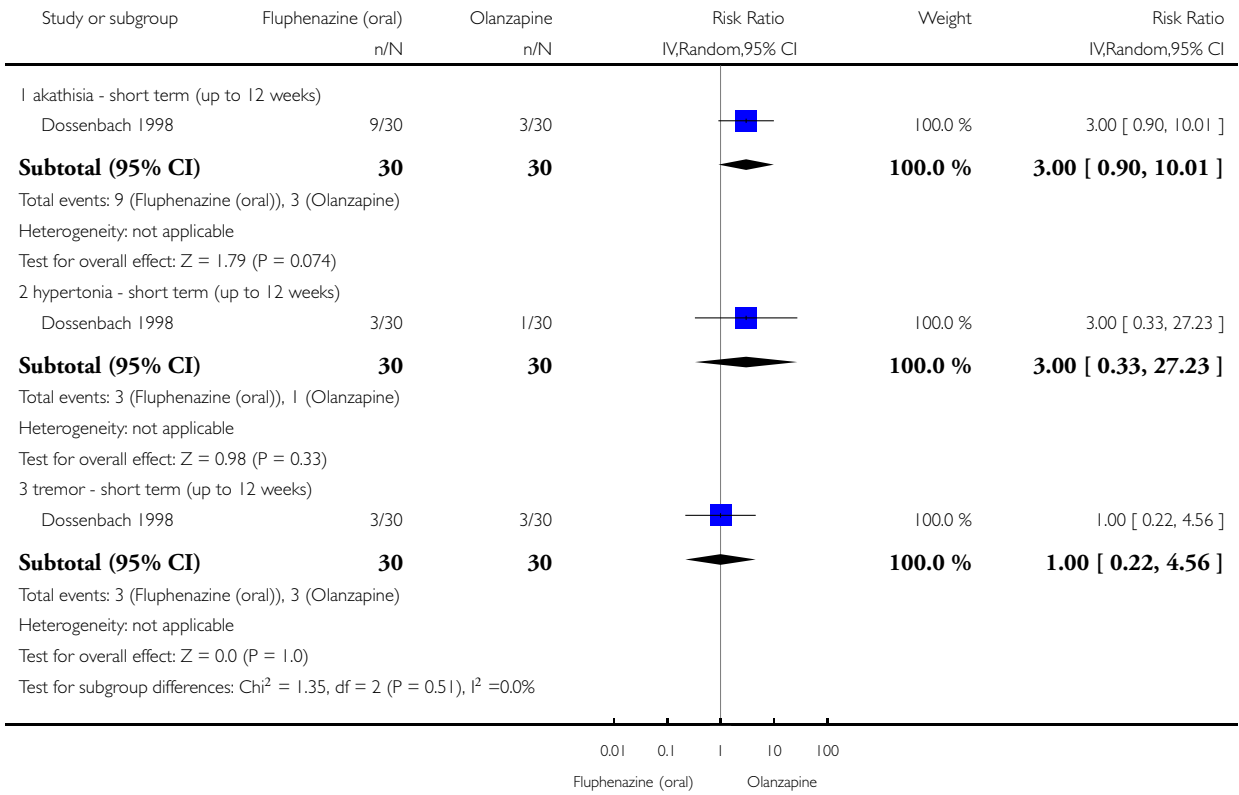


Analysis 4.16. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 16 Adverse effects: 4a. Extrapyramidal effects.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 16 Adverse effects: 4a. Extrapyramidal effects

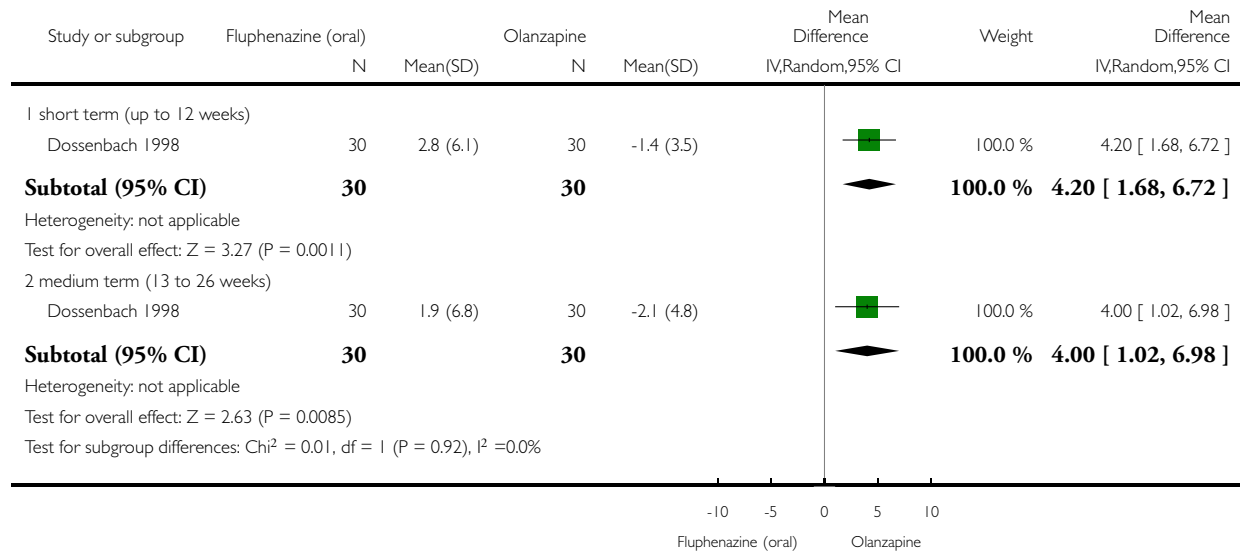


Analysis 4.17. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 17 Adverse effects: 4b. Extrapyramidal effects - average change score (SAS, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 17 Adverse effects: 4b. Extrapyramidal effects - average change score (SAS, high = poor)

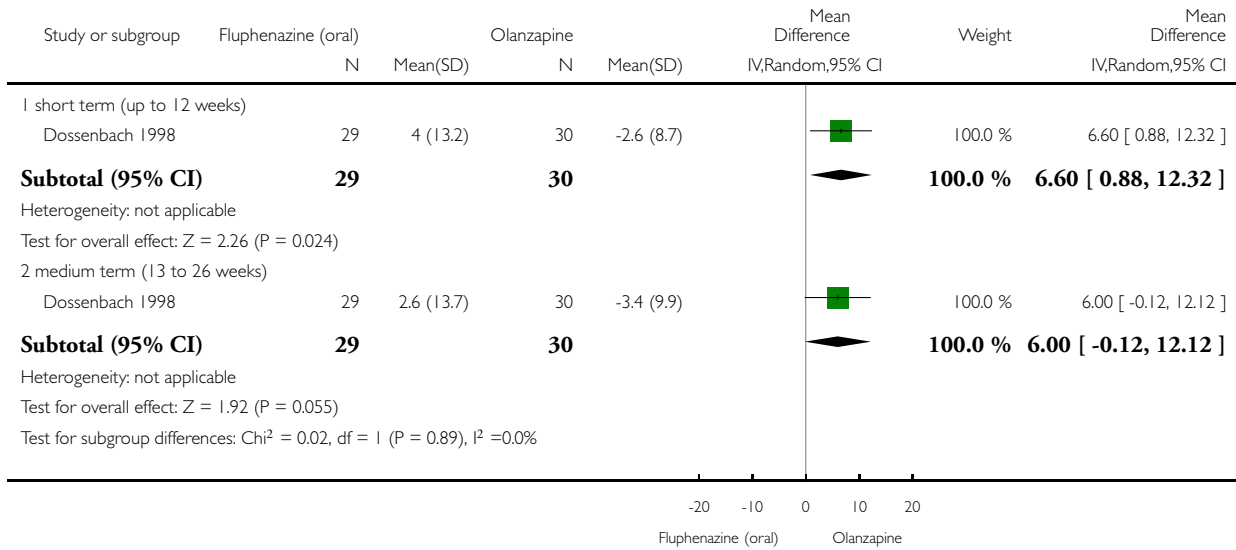


Analysis 4.18. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 18 Adverse effects: 4c. Extrapyramidal effects - average change score (HAS, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 18 Adverse effects: 4c. Extrapyramidal effects - average change score (HAS, high = poor)

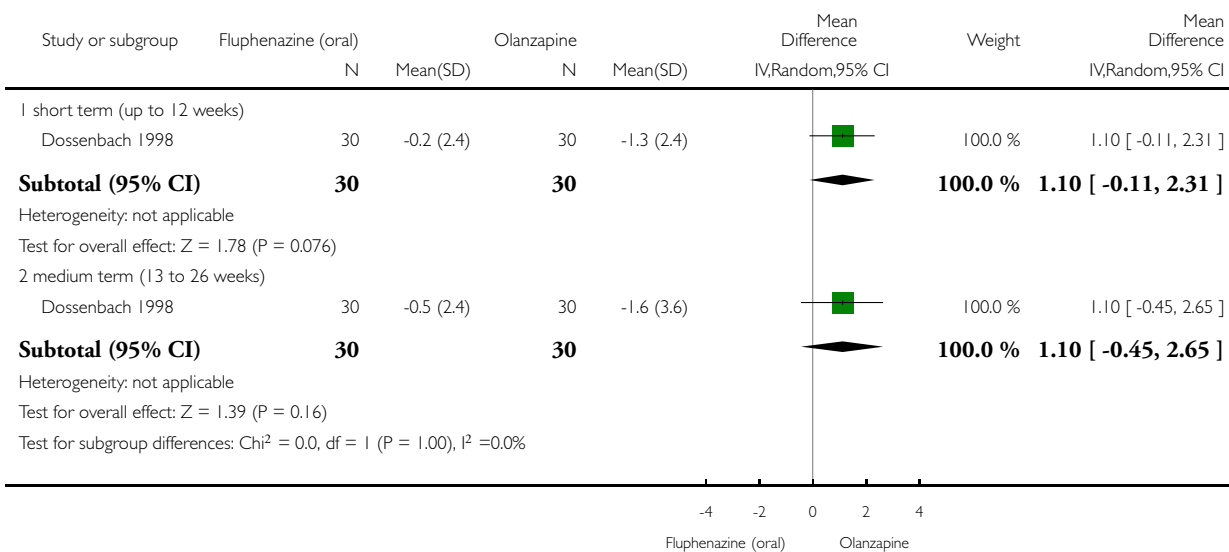


Analysis 4.19. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 19 Adverse effects: 4d. Extrapyramidal effects - average change score (AIMS, high = poor).

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 19 Adverse effects: 4d. Extrapyramidal effects - average change score (AIMS, high = poor)

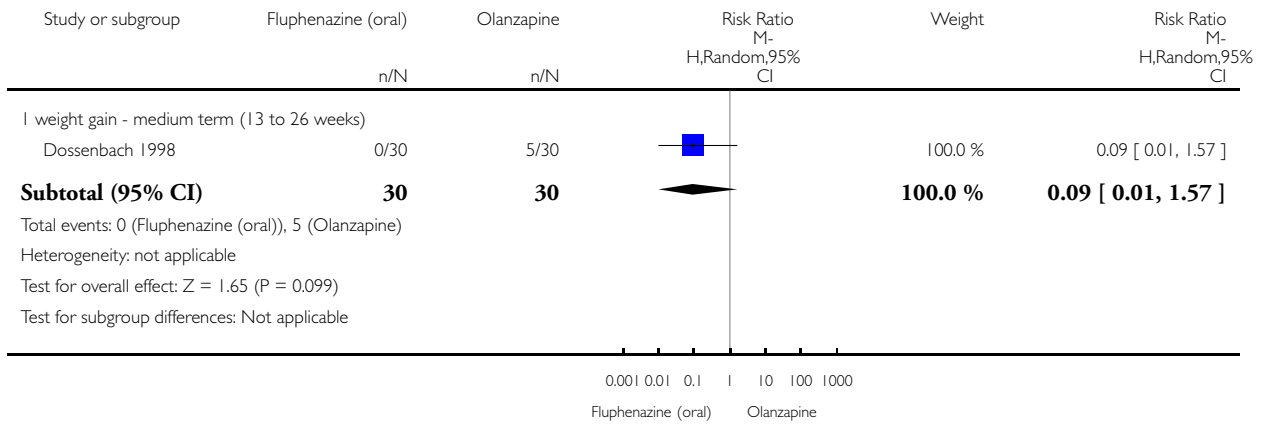


Analysis 4.20. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 20 Adverse effects: 5. Other adverse events.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 20 Adverse effects: 5. Other adverse events

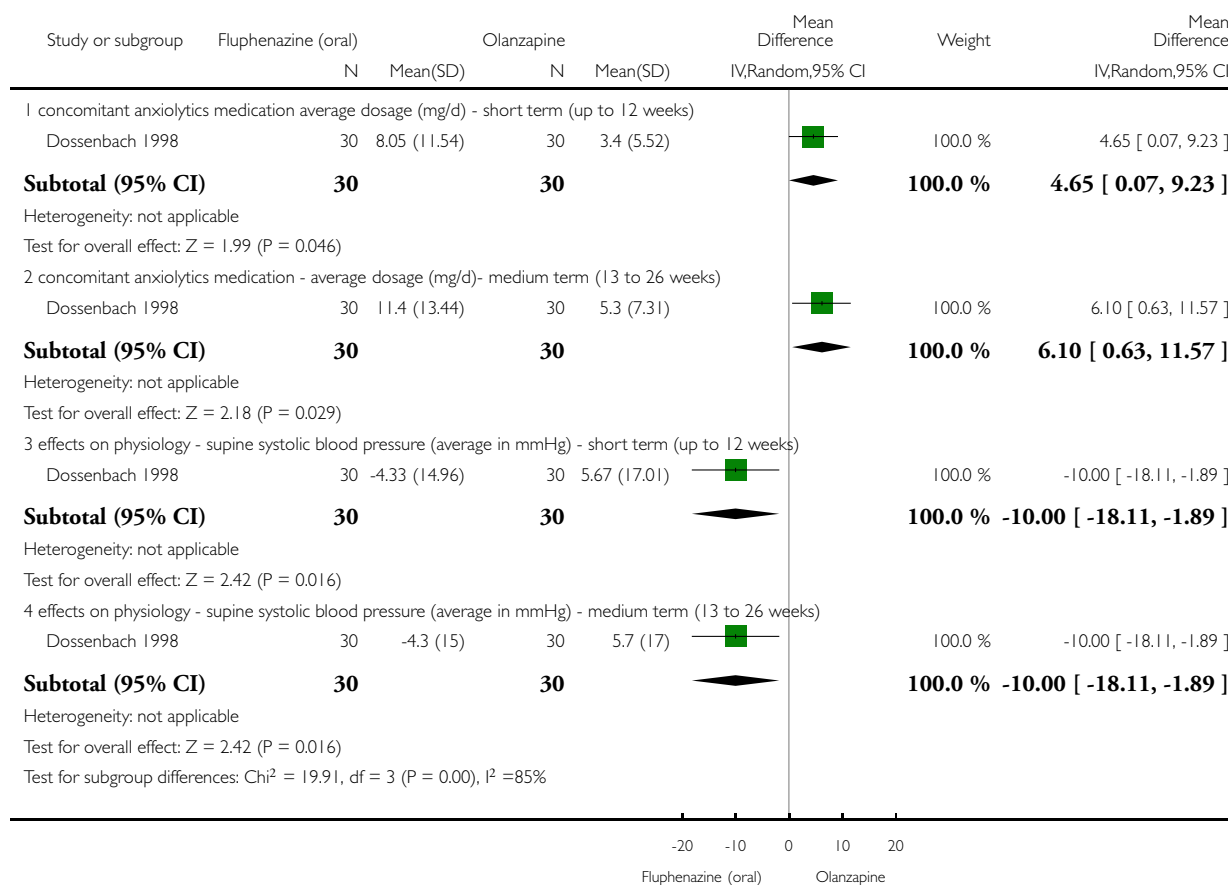


Analysis 4.21. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 21 Adverse effects: 5b. Other adverse events.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 21 Adverse effects: 5b. Other adverse events



Analysis 4.22. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 22 Adverse effects: 5c. Other (skewed).

Adverse effects: 5c. Other (skewed)

Study	Intervention	Mean(kg)	SD	N	weight gain (average weight in kg) - short term (up to 12 weeks)
Dossenbach 1998	Fluphenazine	0.45	2.72	30	weight gain (average weight in kg) - short term (up to 12 weeks)
Dossenbach 1998	Olanzapine	2.43	3.83	30	

Adverse effects: 5c. Other (skewed) (Continued)

weight gain (average weight in kg) - medium term (13 to 26 weeks)					weight gain (av
Dossenbach 1998	Fluphenazine	0.04	3.21	30	
Dossenbach 1998	Olanzapine	5.15	6.41	30	

Analysis 4.23. Comparison 4 FLUPHENAZINE (ORAL) vs OLANZAPINE, Outcome 23 Leaving the study early.

Review: Fluphenazine (oral) versus atypical antipsychotics for schizophrenia

Comparison: 4 FLUPHENAZINE (ORAL) vs OLANZAPINE

Outcome: 23 Leaving the study early

