Supplementary Online Content

Leucht S, Bauer S, Siafis S, et al. Examination of dosing of antipsychotic drugs for relapse prevention in patients with stable schizophrenia: a meta-analysis. *JAMA Psychiatry*. Published online August 18, 2021. doi:10.1001/jamapsychiatry.2021.2130

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This supplementary material has been provided by the authors to give readers additional information about their work.

eAppendix 1. Dose response meta-analysis of antipsychotic drugs for relapse prevention in schizophrenia

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Review question

Antipsychotic drugs are the mainstay of treatment of schizophrenia. The antipsychotic doses which are effective for the acute treatment of the disorder have been relatively well established (Leucht et al. American Journal of Psychiatry 2019;177:342-53). It is, however, unclear whether lower doses are sufficient for maintenance treatment (relapse prevention) than in the acute phase. To know this would be important due reduce the side-effects of antipsychotics to a minimum. We aim to fill this gap by dose-response meta-analysis of randomized controlled trials. Two separate publications are planned. One on efficacy with the primary outcome relapse (study defined) and one on the major side-effects of antipsychotics drugs, i.e. 1) at least one side-effect (primary outcome), 2) weight gain, 3) extrapyramidal side-effects, 4) prolactin increase, 5) QTc prolongation, 6) sedation.

Searches

- Electronic databases: We will search the Cochrane Schizophrenia Group's Study-Based Register of Trials. The search strategy is: (*Amisulpride Dosage* OR *Aripiprazole Dosage* OR *Asenapine Dosage* OR *Brexpiprazole Dosage* OR *Cariprazine Dosage* OR *Clozapine Dosage* OR *Haloperidol Decanoate Dosage* OR *Haloperidol Dosage* OR *Iloperidone Dosage* OR *Lumateperone Dosage* OR *Lurasidone Dosage* OR *Olanzapine Dosage* OR *Paliperidone Dosage* OR *Paliperidone Palmitate Dosage* OR *Quetiapine Dosage* OR *Risperidone Dosage* OR *Sertindole Dosage* OR *Ziprasidone Dosage* OR *Zotepine Dosage*) in Pairwise Comparison Field of Study Records. This register is compiled of regular searches in multiple electronic databases, clinicaltrials.gov, WHO register of clinical trials and more. Details on the register can be found in (Shokraneh and Adams Systematic Reviews 2019;8:129, Shokraneh and Adams BioImpacts : BI 2017;7:209-17, Health Information and Libraries Journal 2020, Schizophrenia Bulletin Open 2020).
- 2. Previous reviews: We will search the studies of a Cochrane review on the effects of antipsychotic drugs for maintenance treatment in general (Leucht et al. Cochrane Database Syst Rev 2012;Cd008016, Ceraso et al. Cochrane Database Syst Rev 2020). For this review exhaustive searches had been undertaken.
- 3. Reference searching: Reference lists of newly included records will be hand-searched for potentially relevant studies.
- 4. We will contact authors or pharmaceutical companies for missing data of studies published from 1990 onward as long as e-mail addresses were available.

There will be no date/time, language, document type, and publication status limitations. All publications will be selected independently by at least two reviewers. In case of doubt, a third reviewer (SL and JST) will be involved. If this procedure does not lead to resolution of the issue, the study authors will be contacted.

Search strategy

Types of study to be included

- Randomized controlled trials (RCTs) with at least one fixed, antipsychotic dose versus a placebo (active or inactive, e.g. a benzodiazepine) or RCTs which compared at least two fixed doses of the same antipsychotic will be included.
- Both open and blinded trials.
- In case of crossover trials, only data from the first of cross-over phase will be used in order to avoid carry-over effects (Elbourne et al. Int J Epidemiol 2002;31:140-9)

- Cluster randomized trials will be excluded due to the unit-of-analysis-problems associated with this design (Whiting-O'Keefe et al. Med Care 1984;22:1101-14).
- Studies with a high risk of bias in terms of randomization according the Cochrane risk of bias tool will be excluded.
- There will be no language restriction. Studies from mainland China will be excluded due to frequent quality problems (Tong et al. BMC Med Res Methodol 2018;18:96). The reports are usually short making it impossible to detect these problems and authors often do not reply to requests in our experience. Studies conducted in China by international companies will be accepted.
- The minimum study duration will at least 14 weeks. The rational of this cutoff is to exclude short-term trials which usually examine acutely ill patients with schizophrenia. The purpose of the cutoff therefore is to reduce clinical and methodological heterogeneity. It corresponds to the category for short-term trials of the Cochrane Schizophrenia Group (<u>https://schizophrenia.cochrane.org</u>). There will be no a priori defined maximum duration, although we expect that few studies will last longer than 1 year and that the longest trial duration will be 3 years (Ceraso et al. Cochrane Database Syst Rev 2020).
- We will include both, studies which randomize participants in their maintenance phase, and socalled continuation studies, as long as all acute phase responders could be followed up. Continuation are studies in which patients are randomized in the acute phase, and the responders in the acute phase are then followed up and examined for relapse prevention. Designs that allow that participants switch from one randomised group to the other will be excluded. This kind of trials is, for example, often used in cost effectiveness studies, but it is not appropriate to compare the effects of different doses of one drug.
- Studies on the acute treatment of schizophrenia will be excluded.

Condition or domain being studied

Schizophrenia and schizophrenia-related disorder

Participants/population

- Participants with a diagnosis of schizophrenia or schizophrenia-related disorders, e.g. schizophreniform or schizoaffective disorders. We will accept both clinical diagnosis and diagnosis based on operationalized diagnostic criteria. Studies including participants with a diagnosis other than schizophrenia-related disorders will be accepted when the at least 80% of the participants had diagnoses with schizophrenia-related disorders.
- Participants must be in the stable phase of their illness, we will exclude studies in acutely ill patients. Any definition of "stability" will be accepted, because no uniform definition is available.
- Studies in children and adolescents, in elderly patients, in participants with predominant negative symptoms and in participants with a first-episode of schizophrenia will be analyzed separately, because there is evidence that such patients need lower doses, at least in the acute phase (Oosthuizen et al. Int J Neuropsychopharmacol 2004;7:125-31, Krause et al. Eur Neuropsychopharmacol 2018;28:659-74, Krause et al. Eur Arch Psychiatry Clin Neurosci 2018;268:625-39, Krause et al. Eur Neuropsychopharmacol 2018;28:1360-70). It is planned to analyze these populations in separate publications. Studies in treatment resistant patients and in patients with concomitant substance abuse will be analyzed together with the studies on general adults with schizophrenia, but they will be excluded in a sensitivity analysis (Krause et al. Eur Neuropsychopharmacol 2019;29:32-45).
- There will be no other restriction in terms of setting, gender, nationality and ethnicity.

Intervention(s), exposure(s)

 Any of the following antipsychotic drugs will be eligible: amisulpride, aripiprazole (oral, depot formulations of maintena and lauroxil), asenapine (oral and transdermal), brexpiprazole, cariprazine, clozapine, haloperidol (oral and depot), fluphenazine, iloperidone, lumateperone, lurasidone, olanzapine (oral and depot), quetiapine, paliperidone (oral and depot), risperidone (oral, depot formulations of consta and RBP-7000), sertindole, ziprasidone, zotepine. This selection comprises all so-called second-generation antipsychotic drugs available in Europe and/or the US. Haloperidol will also be examined, because it was the gold standard in many countries before these more recent drugs had been developed. As the depot formulation of fluphenazine was the best investigated drug for dose-response before the advent of the second-generation antipsychotics, we included this other standard first-generation antipsychotic, as well.

- There will be no restriction in terms of route of administration (except for short-acting injections and intranasal forms that are used for acute agitation). Antipsychotic compounds given via different route of administration will be considered as separate compounds. For example, oral aripiprazole, aripiprazole maintena and aripiprazole lauroxil will be considered as three separate antipsychotic interventions. In a similar vein for oral asenapine/transdermal asenapine, oral paliperidone/paliperidone depot once monthly, risperidone/risperidone consta/risperidone RBP-7000, olanzapine/olanzapine depot.
- Fixed-dose schedules, and studies in which patients are randomised to different, narrow, nonoverlapping, fixed dose range, for example olanzapine 5mg/day +/- 2.5 mg/day versus olanzapine 10mg/day +/-2.5mg/day. Flexible-dosing schedules will not be eligible.

Comparator(s)/control

Placebo, active of non-active.

Context

We will include studies in outpatient and in inpatient settings as long as the patients are stable at baseline. We will not generally exclude inpatient studies, because there might be studies which have been conducted in long-term wards for stable patients. The detailed in- and exclusion criteria are listed above.

Main outcome(s)

We plan two separate publications, one with a focus on efficacy, the other one on different side-effects. We wrote one protocol, because the overall methodology and the searches will be the same. The following are the outcomes for the efficacy focused review.

The number of participants relapsed (study defined)

We will accept any definition of relapse. Different definitions have been used in the literature and that there is no consensus as to which is the most appropriate one. We will, however, prefer relapse criteria which are operationalized by rating scales rather than other criteria that we will extract following this hierarchy a) 'patient relapsed by the judgment of the clinician/rater', b) need for additional antipsychotic medication c) dropout due to inefficacy d) re-hospitalisation and e) and other.

The primary outcome relapse will be extracted at 6 months, 9 months, 12 months and longer than 12 months. We will statistically analyze the outcome closest to 12 months. All other outcomes will be measured at study endpoint.

* Measures of effect

Relapse will be analyzed with odds ratios. Also see section 'strategy for data synthesis'.

Additional outcome(s)

The following secondary outcomes will be analysed in the review.

- 1. Mean change from baseline to endpoint of overall symptoms of schizophrenia as measured by the Positive and Negative Syndrome Scale (Kay et al. Schizophr Bull 1987;13:261-76), the Brief Psychiatric Rating Scale (Overall and Gorham Psychological Reports 1962;10:799-812) or any other published scale to measure the symptoms of schizophrenia.
- 2. Premature study discontinuation due to any reason. This outcome is actually a measure of effectiveness, because it comprises dropouts due to side-effects, inefficacy and others.
- 3. Rehospitalisation for psychiatric reasons
- 4. Premature study discontinuation due to adverse events, where dropouts due to side-effects were preferred whenever available.

* Measures of effect

Rating scales of schizophrenia symptoms will be analyzed with the standardized mean difference, because we expect that different scales have been used in the studies. All other outcomes are dichotomous for which odds ratios will be use as measures of effect. Also see section 'strategy for data synthesis'.

Data extraction (selection and coding)

1. Selection of trials: At least two reviewers will independently inspect the titles and abstracts of non-duplicated references identified through the search and will exclude those not pertinent. Discrepancies between the two reviewers will be resolved by discussion. If doubts still remain, the full text will be obtained and eligibility will be assessed. Full texts of included references will be obtained and independently assessed by two reviewers for eligibility. Again, disagreements will be resolved by discussion and, if needed, a third author will be involved (SL or JST). When required, further information will be requested from study authors.

2. Data extraction: Two authors will independently extract data from all selected trials in a Microsoft Access database. When disagreement arises, we will resolve it by discussion and, if needed, involving a third senior author (SL or JST). Where this is not sufficient, we will contact the study authors.

- When authors of original studies used imputation methods to handle missing data, we will prefer them to completers' data. For the outcome relapse we will prefer data based on survival analysis rather than the absolute number of participants relapsed. In terms of continuous data mixed-models of repeated measurement (MMRM), multiple imputation will be preferred over last-observation carried forward (LOCF), if available.
- For dichotomous outcomes, if only completer analyses are presented, we will assume that participants lost to follow-up did not have the outcome. We think that another assumption would overestimate the risk.
- For continuous outcomes, we will prefer change scores to follow-up data, but we will also accept the latter when the former are not available.
- Missing SDs will be calculated from 1) standard error (SE), 2) other measures of variability (95% confidence intervals, ranges etc), 3) test statistics 4) imputed from the SDs of the other studies using a validated method (Furukawa et al. J Clin Epidemiol 2006;59:7-10) according to the Cochrane Handbook (Higgins and Green 2011;4).

Risk of bias (quality) assessment

Two independent review authors will assess the risk of bias in the selected studies using the 'Cochrane Collaboration risk of bias' tool version 2 (Sterne et al. Bmj 2019;366:I4898). When disagreement arises we will resolve it by discussion and, if needed, involving a third senior author.

Strategy for data synthesis

In the efficacy publication we will – in addition to analyzing each antipsychotic separately - pool all studies after converting the doses to risperidone equivalents based on the following criteria: Dose equivalence based on 95% Effective Doses (Leucht et al. American Journal of Psychiatry 2019;177:342-53), if not available Minimum Effective Dose method (Leucht et al. Schizophr Bull 2014;40:314-26, Rothe et al. Schizophr Res 2018;193:23-8), if not available Mean Dose method (Leucht et al. Schizophr Bull 2015;41:1397-402, Davis J Psychiatr Res 1974;11:65-9), if not available Daily Defined Dose method (Leucht et al. Schizophr Bull 2016;42 Suppl 1:S90-4) if not available based on the International Consensus of Antipsychotic Doses (David M. Gardner et al. American Journal of Psychiatry 2010;167:686-93). In a secondary analysis we will convert doses based on the expert opinions according to the International Consensus of Antipsychotic Doses (David M. Gardner et al. American Journal of Psychiatry 2010;167:686-93) supplemented by similar judgements by the reviewer team for drugs that were not reported in the consensus statement.

- The effect sizes for dichotomous outcomes will be the odds ratio (OR). The effect sizes for continuous rating scales for efficacy we will use the standardized mean difference (SMD as Hedges' g), because we expect that various scales have been used in the studies to measure the same concepts. All effect sizes will be accompanied by their 95% confidence intervals.
- We will conduct a one-stage dose response meta-analysis in a frequentist framework using restricted-cubic splines with the R package 'dosresmeta' developed by Crippa et al (Crippa et al. Stat Methods Med Res 2019;28:1579-96, Crippa and Orsini Journal of Statistical Software 2016;72). We will use knot points at the 25th, 50th and 75th percentile.
- We will produce absolute dose-response curves: we will synthesize the effects in the placebo arms and we will transform the relative dose-response curves estimated in previous steps to absolute curves.
- For drugs with enough data we will use the Wald statistic to explore whether there is evidence of an overall dose-response relationship and we will report the p-values.
- We will use the dose-response curves to estimate the 95% effective dose (ED95) and 50% effective dose (ED50) as is customary in dose-response analysis (1, 4). The ED50 is the mean dose that produces 50% of the maximum relapse prevention compared with placebo, and the ED95 is the mean dose that produces 95% of the maximum reduction. The ED50 and the ED95 will be calculated for each drug separately and for all drugs pooled.
- Small study effects and the possibility of publication bias will be assessed with funnel plots and Egger's test, when there are at least 10 studies available.

Analysis of subgroups or subsets

Predefined sensitivity analyses of the primary outcomes will be:

- In the primary analysis, we will synthesize studies that compared at least two doses of a compound with placebo. In a sensitivity analysis we will exclude studies that compared only a *single* dose of an antipsychotic with placebo, because such studies were not designed to address dose-response and could therefore produce methodological heterogeneity.
- Immediate (IR) and extended release (XR) formulations will be analyzed separately (i.e. for quetiapine).
- We will exclude open RCTs for subjective outcomes.
- We will examine the subgroup of patients in remitted (added post-hoc).

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Leucht S, Samara M, Heres S, Patel MX, Woods SW, Davis JM. Dose equivalents for second-generation antipsychotics: the minimum effective dose method. Schizophr Bull. 2014;40:314-26.
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Changes to the original protocol published in Prospero

- This version refers to the efficacy focussed review, additional ones on specific side-effects are planned. This decision was made a priori, see first paragraph of the protocol.

- We added a subgroup analysis of patients in remission at baseline. The single first-episode study was analysed together with the studies in chronic patients, but excluded in a sensitivity analysis. Similarly, due to a relative scarcity of data we pooled oral and depot formulations, but we also analysed both separately (data not shown).

95% and 50% effective doses were not calculated.

- The outcome rehospitalisation for psychiatric reasons was analysed separately in addition to its analysis as an indicator for relapse.

- Following reviewer requests we tested with linear splines up to which dose the dose-response curve still showed a significantly increasing slope, and added subgroup analyses on long-acting injectable versus oral medications, first-generation versus second-generation antipsychotics, percentage male and age.

eAppendix 2. . Description of the search strategy

1 Database

Cochrane Schizophrenia Group's Study-Based Register of Trials. Details are also described on the Cochrane Schizophrenia Group's website https://schizophrenia.cochrane.org/.

2 Date of search

9th March 2020

3 Search Strategy

Strategy: (*Amisulpride Dosage* OR *Aripiprazole Dosage* OR *Asenapine Dosage* OR *Brexpiprazole Dosage* OR *Cariprazine Dosage* OR *Clozapine Dosage* OR *Fluphenazine Dosage* OR *Haloperidol Decanoate Dosage* OR *Haloperidol Dosage* OR *Iloperidone Dosage* OR *Lumateperone Dosage* OR *Lurasidone Dosage* OR *Olanzapine Dosage* OR *Paliperidone Dosage* OR *Paliperidone Palmitate Dosage* OR *Quetiapine Dosage* OR *Risperidone Dosage* OR *Sertindole Dosage* OR *Ziprasidone Dosage* OR *Zotepine Dosage*) in Pairwise Comparison Field of Study Records

4 Search Results

There were 1306 references from 390 studies.

5 References to database details

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3. Shokraneh, Farhad; Adams, Clive E. Study-based registers reduce waste in systematic reviewing: discussion and case report. *Systematic Reviews* 2019; 8: 129. DOI 10.1186/s13643-019-1035-3

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5 Update search in Pubmed January 1st 2021

Search: (amisulpride OR aripiprazole OR asenapine OR brexpiprazole OR cariprazine OR clozapine OR fluphenazine OR haloperidol* OR iloperidone OR lumateperone OR lurasidone OR olanzapine OR paliperidone OR quetiapine OR risperidone OR sertindole OR ziprasidone OR zotepine) Filters: Randomized Controlled Trial, From 2020/1/1 to 2021/1/1

28 selected items

eFigure. PRISMA diagram of the search



N = number of studies, * some publications provided data on two studies

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Study	Year of Publi catio nm	Degree of blinding	Dur atio n (wk s)	Diagnostic term (diagnostic criteria)	Interventions	Appli catio n	Dosing interval	Dose mean and range mg	Number of patient s random ized	Mea n age in year s
					Placebo	oral	-	-	71	48.7
Arato et al	2002	double	52	Schizophrenia	Ziprasidone 160mg	oral	daily	160 (160-160)	67	49.6
2002(1)		biind	(DSM-III-R)	(DSM-III-R)	Ziprasidone 40mg	oral	daily	40 (40-40)	72	50.8
				Ziprasidone 80mg	oral	daily	80 (80-80)	68	49.8	
Velligan et al					Haloperidol	oral	daily	12 (12-12)	41	37
2002(2)	1005	double	50	Schizophrenia (DSM-III-R)	Quetiapine 300mg	oral	daily	300 (300-300)	88	37
(unpublished	1995	blind	52		Quetiapine 600mg	oral	daily	600 (600-600)	87	38
data obtained)					Quetiapine 75mg	oral	daily	75 (75-75)	85	39
5					Haloperidol 15±5mg	oral	daily	15.8 (10-20)	10	36.1
Beasley et al 1996a (Study				.	Olanzapine 10±2.5mg	oral	daily	11.6 (7.5- 12.5)	9	40.9
F1M-MC- HGAD)(3)	1996	double blind	46	Schizophrenia (DSM-III-R)	Olanzapine 15±2.5mg	oral	daily	15.8 (12.5- 17.5)	22	34.4
(unpublished data obtained)	ished tained)			Olanzapine 5±2.5mg	oral	daily	6.3 (2.5-7.5)	14	33.5	
					Placebo	oral	-	-	13	36.1
Beasley et al 1997 (Study	1997	double blind	46	Schizophrenia (DSM-III-R)	Haloperidol 15±5mg	oral	daily		28	34.9

eTable1: Characteristics of Included Studies

F1D-EW- E003)(3)					Olanzapine 1±2.5mg	oral	daily	1 (1-1)	33	34.0
(unpublished data obtained)					Olanzapine 10±2.5mg	oral	daily	11.3 (7.5- 12.5)	40	36.5
					Olanzapine 15±5mg	oral	daily	16.1(12.5- 17.5)	43	37.6
					Olanzapine 5±2.5mg	oral	daily	6.6 (2.5-7.5)	31	35.2
Carpenter et al	1000	double	E A	Schizophrenia or	Fluphenazine 25mg/2 weeks	depot	every 2 weeks	25 (25-25)	25	34.7
1999(4)	1999	blind	54	(DSM-III-R)	Fluphenazine 25mg/6 weeks	depot	every 6 weeks	25 (25-25)	25	36. 2
				First-episode	Placebo	oral	-	-	89	24.9
Chen et al 2010(5)#	2010	double blind	52	outpatients with schizophrenia or related disorders (DSM-IV). <u>Remission criteria</u> : no history of relapse in the last 12 months (defined as any increase in positive symptoms leading to adjustement of drug treatment or readmission to hospital, ascertained with information from patients, carers, case managers, clinicians, and clinical records). All were non-psychotic at study entry as defined by having below threshold scores on five	Quetiapine	oral	daily	400 (400-400)	89	23.5

				key psychotic symtpoms on the PANSS: P1 delusions ≤2, P2 conceptual disorganization ≤ 3, P3 hallucinatory behavior ≤ 2, P6 suspiciousness ≤ 4, G9 Unusual thought content ≤3; and ≤2 (borderline or questionable illness) on the CGI severity.						
Cooper et al	2000	double	26	Schizophrenia	Placebo	oral	-	-	58	41.6
2000(6)		biina		(DSM-III-R)	Zotepine	oral	daily	n.i. (150-300)	63	43
Dotti et al	1979	double	39	Schizophrenia (Clinical	Fluphenazine	depot	every 4 weeks	n.i. (25-50)	10	n.i.
1979(7)		blind		diagnosis)	Placebo	depot	-	-	10	n.i.
Eklund et al	1991	double	48	Schizophrenia (Research Diagnostic	Haloperidol	depot	every 4 weeks	60 (60-60)	20	51.7
1991(8)		blind		Criteria (RDC))	Placebo	depot	-	-	23	51.7
Fleischhacker et	2014	double	20	Schizophrenia	Aripiprazole LAI 400mg/4weeks	depot	every 4 weeks	392 (300-400)	265	41.7
al 2014(9)	2014	blind	38	(DSM-IV-TR)	Aripiprazole LAI 50mg/4weeks	depot	every 4 weeks	50 (25-50)	131	40.2
					Paliperidone 100mg	depot	every 4 weeks	100 (100-100)	86	43
Hough et al 2009(10)	2009	double blind	25	Schizophrenia (DSM-IV)	Paliperidone 50mg	depot	every 4 weeks	50 (50-50)	82	43
					Paliperidone 75mg	depot	every 4 weeks	75 (75-75)	84	43
Huttunen et al 1996(11)	1996	open-label	104	Schizophrenia (DSM-III-R)	Haloperidol 150mg	depot	every 4 weeks	150 (150-150)	13	n.i.

					Haloperidol 25mg	depot	every 4 weeks	25 (25-25)	13	n.i.
				Schizophrenia, any subtype (RDC) in good	Fluphenazine	depot	every 2 weeks	n.i. (1.25-5)	8	26.5
Kane et al 1979(12)#	1979	double blind	26	remission with no evidence for significant psychopathology and with reasonably good social and vocational functioning for at least 6 months during a prospective open non- randomised trial with fluphenazine decanoate. All patients were being successfully maintained in the community.	Placebo	depot	-	-	8	26.9
				Outpatients with schizophrenia or	Fluphenazine 1.25-5mg	depot	every 2 weeks	3.125 (1.25-5)	66	28.9
Kane et al 1983(13)#	1983	double blind	52	schizoaffective disorder (RDC) in <u>remission or</u> <u>partial remission</u> defined by GAS \geq 35; BPRS \leq 4 on conceptual disorganization and hallucinatory behavior, \leq 5 on suspiciousness, and \leq 3 on unusual thought content. No fluctuation in either direction by more than 10 on the GAS and by more than 1 on any of the BPRS items	Fluphenazine 12.5-50mg	depot	every 2 weeks	31.25 (12.5- 50)	66	28.9

				mentioned. Patients were required to maintain this stability while receiving a constant IM dose of fluphenazine-decanoate within the range of 12.5 to 50mg/2week. Average number of weeks from hospital discharge to study entry 63.9+-78.5 weeks.						
				Schizophrenia or schizoaffective disorder (DSM-III) outpatients	Haloperidol 100mg Haloperidol	depot depot	every 4 weeks every 4	100 (100-100) 200 (200-200)	31 29	38 40
				with a baseline state of relative <u>remission</u> for at least 3 months during maintenance treatment with antipsychotic medication and a BPRS ≤3 on conceptual disorganization and unusual thought content and ≤4 on hallucinatory behavior and suspiciousness	Haloperidol 25mg	depot	every 4	25 (25-25)	29	38
Kane et al 2002b(14)#	2002	double blind	52		Haloperidol 50mg	depot	every 4 weeks	50 (50-50)	30	38
Kane et al	2010	double	24	Schizophrenia	Olanzapine LAI 150mg/2weeks	depot	every 2 weeks	150 (150-150)	140	37.7
2010(15)	2010	blind	24	(DSM-IV or DSM-IV-TR)	Olanzapine LAI 300mg/2weeks	depot	every 2 weeks	300 (300-300)	141	39.5

					Olanzapine LAI 405mg/4weeks	depot	every 4 weeks	405 (405-405)	318	39
					Olanzapine LAI 45mg/4weeks	#NV	n.i.	n.i. (n.in.i.)	#NV	n.i.
Kane et al	2012	double	52	Schizophrenia	Aripiprazole	depot	every 4 weeks	396.3 (300- 400)	269	40.1
2012(16)		blind		(DSM-IV-TR)	Placebo	depot	-	-	134	41.7
					Aripiprazole 200mg	depot	every 4 weeks	200 (200-200)	11	46
Mallikaarjun et al 2013(17)	2013	open-label	24	Schizophrenia (DSM-IV-TR)	Aripiprazole 300mg	depot	every 4 weeks	300 (300-300)	16	43.3
					Aripiprazole 400mg	depot	every 4 weeks	400 (400-400)	14	46.8
Marder et al	1084	double	104	Schizophrenia (DSM-III)	Fluphenazine 25mg	depot	every 2 weeks	25 (25-25)	31	38.1
1984(18)	1304	blind	104		Fluphenazine 5mg	depot	every 2 weeks	5 (5-5)	35	35.1
McEvoy et al 2007 extension		double		Schizophrenia	Aripiprazole 10- 15mg	oral	daily	n.i. (10-15)	88	n.i.
(19) (unpublished data obtained)	2005	blind	46	(DSM-IV)	Aripiprazole 20- 30mg	oral	daily	n.i. (20-30)	88	n.i.
				Fully <u>remitted</u>	Haloperidol	oral	daily	3 (3-3)	10	33
Nishikawa et al 1982(20)#	1982	double blind	156	schizophrenic outpatients who were working in society	Placebo	oral	-	-	10	33.6
				Symptom free	Haloperidol 1mg	oral	daily	1 (1-1)	13	40
				outpatients	Haloperidol 3mg	oral	daily	3 (3-3)	12	36.3
Nishikawa et al	1984	double	52	(DSM-III) in the	Haloperidol 6mg	oral	daily	6 (6-6)	12	42.8
1984(21)#		blind		recovery stage of remission or residual phase	Placebo	oral	-	-	13	38.8
	2016		28	Schizophrenia	Lurasidone	oral	daily	78.9 (40-80)	144	43

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Tandon et al 2016(22)		double blind		(DSM-IV-TR)	Placebo	oral	-	-	141	42.4
Pigott et al	2002	double	26	Schizophrenia	Aripiprazole	oral	daily	15 (15-15)	155	42.2
2003(23)	2003	blind	20	(DSM-IV)	Placebo	oral	-	-	155	41.7
				Schizophrenia (any subtype), schizoaffective	Fluphenazine 12.5-50mg	depot	every 2 weeks	27.47353 (12.5-50)	55	29.6
1997(24)	1997	blind	104	disorder or schizophreniform	Fluphenazine 2.5- 10mg	depot	every 2 weeks	13.775 (2.5- 10)	52	29.6
				(DSM-III-R)	Placebo	depot	-	-	49	29.6
Simpson et al		double		Schizophrenia or	Risperidone 25mg	depot	every 2 weeks	25 (25-25)	163	41.7
2006(25)	2006	blind	52	(DSM-IV)	Risperidone 50mg	depot	every 2 weeks	50 (50-50)	161	40.2

n.i. = not indicated, DSM = Diagnostic and Statistical Manual, RDC = Research diagnostic criteria , # studies in remitted patients.

eTable2: Characteristics of Excluded Studies

Author	Reason for exclusion
Agid et al 2007(26)	Method: short-term
Agid et al 2008(27)	Method: short-term
Anderson et al 1976(28)	Method: short-term
Andrews et al 1976(29) ⁱ	Method: randomized
	Participants: schizophrenia
	Intevention: flexible dose ranges
Anonymus 1994(30)	Method: short-term
Anonymus 2001(31)	Method: short-term
Aravagiri et al 1994(32)	Method: randomized
	Participants: schizophrenia
	Intevention: fixed dose
	Outcome: not relapse prevention
Arvanitis et al 1997(33)	Method: short-term
Assion et al 2008(34)	Method: short-term
Auby et al 2002(35)	Method: short-term
Bark et al 1996(36)	Method: short-term
Barnas et al 2001(37)	Method: short-term
Bateman et al 1979(38)	Method: short-term
Beasley et al 1996 (Study F1D-MC-	Method: randomized
HGAP)(39)	Participants: schizophrenia
	Intervention: no fixed doses
Beasley et al 2003(40)	Method: randomized
	Participants: schizophrenia
	Intevention: flexible dose range
Berger et al 2008(41)	Method: short-term
Berwaerts et al 2015(42)	Method: randomized
	Participants: schizophrenia
	Intevention: not all participants received the
	same dosages of paliperidone palimtate
Bitter et al 1989(43)	Method: short-term
Bjorndal et al 1980(44)	Method: short-term
Boyer et al 1987(45)	Method: short-term
Brambilla et al 1987(46)	Method: randomized
	Participants: chronic schizophrenia
	Intevention: combination of drugs
Bristol-Myers 2004/CN138114(47)	Method: short-term
Bristol-Myers 2005/CN138050(48)	Method: short-term
Bunney et al 2010(49)	Method: short-term
Cada et al 2004(50)	Method: short-term
Canuso et al 2010(51)	Method: short-term
Cetin et al 1999(52)	Method: short-term
Chang et al 1991(53)	Method: short-term
Channabasavanna et al 1987(54)	Method: short-term
Chapel et al 2009(55)	Method: short-term
Chavda et al 2004(56)	Method: short-term
Citome et al 2019(57)	Method: short-term
Clark et al 1975(58)	Method: randomized
	Participants: chronic schizophrenia
	Intevention: flexible dosage
Clerc et al 1989(59)	Method: short-term
Coppola et al 2011(60)	Method: short-term

Correll et al 2015(61)	Method: short-term
Correll et al 2020(62)	Method: short-term
Ctri-2014-04-004521(63)	Method: short-term
Cutler et al 2006(64)	Method: short-term
Cutler et al 2010(65)	Method: short-term
Czobor et al 1993(66)	Method: short-term
Daniel et al 1999(67)	Method: short-term
Daniel et al 2001(68)	Method: short-term
Daniel et al 2001a(69)	Method: short-term
Davidson et al 2007(70)	Method: short-term
Davis et al 1985(71)	Method: short-term
De Buck et al 1973(72)	Method: short-term
DelBello et al 2008(73)	Method: short-term
Dencker et al 1978(74)	Method: short-term, cross over design, 4 wks
	each
Denijs et al 1973(75)	Method: randomized
	Population: stable schizophrenics
	Intervention: no fixed dosage
do Carmo Borges 2012(76)	Method: short-term
Donlon et al 1978(77)	Method: short-term
Donlon et al 1980(78)	Method: short-term
Dubitsky et al 2002(79)	Method: randomized
	Population: stable schizophrenia
	Intervention: no fixed dose
Dubovsky et al 2012(80)	Method: short-term
Durgam et al 2014(81)	Method: short-term
Durgam et al 2015(82)	Method: short-term
Durgam et al 2016(83)	Method: short-term
Durgam et al 2016a(84)	Method: randomized
	Population: stable schizophrenia
	Intervention: no fixed dose
Dutoit et al 1995(85)	Method: short-term
Eerdekens et al 2004(86)	Method: short-term
Eli Lilly 2008(87)	Method: not adequately randomized
EMEA 2007(88)	Method: pooled data
Euctr2006-006434-17(89)	Method: short-term
Findling et al 2008(90)	Method: short-term
Findling et al 2015(91)	Method: short-term
Findling et al 2015a(92)	Method: short-term
Fleischhacker et al 2003(93)	Method: randomization issues
Fleischhacker et al 2017(94)	Method: randomized
	Population: stable schizophrenia
	Intervention: no fixed dose
Freeman et al 1962(95)	Method: randomized
	Population: stable schizophrenia
	Intervention: flexible doses
Fu et al 2015(96)	Method: randomized
	Population: schizoaffective disorder
	Intervention: flexible doses
Gallant et al 1974(97)	Method: randomized
	Population: severily III schizophrenics
	Intervention: Tiexible doses of pentiuridol
Gitlin et al 1988(98)	Nethod: short-term cross over phases
Gitlin et al 2001(99)	Method: short-term cross over phases

Goff et al 2013(100)	Method: short-term
Goldman et al 2017(101)	Method: short-term
Good et al 1958(102)	Method: short-term cross over phases
Gopal et al 2009(103)	Method: short-term
Grosset al 1974(104)	Method: randomized
	Population: chronic schizophrenia
	Intervention: wrong medication (pimozide,
	trifluoperazine)
Gutierrez et al 1996(105)	Method: short-term
H. Lundbeck 2009(106)	Method: short-term
Haas et al 2009(107)	Method: short-term
Hale et al 2012(108)	Method: short-term
Hard et al 2017(109)	Method: randomized
	Population: schizophrenia
	Intervention: All groups received the same
	dosage over time but in different divisions
Hard et al 2018(110)	Method: short-term
Harvey et al 2009(111)	Method, Randomized
	Population: schizophrenia
	Intervention: no fixed dose
Harvey et al 2010(112)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed dose in continuitation
	phase
Hershon et al 1972(113)	Method: randomized
	Population: schizophrenia
	Intervention: wrong medication (trifluoperazine)
Higuchi et al 2019(114)	Method: short-term
Higuchi et al 2019a(115)	Method: short-term
Hirsch et al 1973(116)	Method: randomized
	Population: chronic schizophrenia
	Intervention: no fixed dosage
Hirsch et al 1989(117)	Method: randomized
	Population: chronic schizophrenia
	Intervention: no fixed dosage
Hirsch et al 1996(118)	Method: randomization issues
Hirschowitz et al 1997(119)	Method: short-term
Hogarty et al 1974(120)	Method: randomized
	Population: schizophrenia
	Intervention: flexible doses
Hogarty et al 1988(121)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed dosages
Hogarty et al 1995(122)	Method: short-term
Horner et al 2012(123)	Method: short-term
Hough et al 2010(124)	Method: randomized
	Population: schizophrenia
	Intervention: flexible doses
Hough et al 2011(125)	Method: short-term
Huber et al 1971(126)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed dosages
Inderbitzin et al 1994(127)	Method: randomized
	Population: schizophrenia

	Intervention: no fixed doses
Ishigooka et al 2018(128)	Method: short-term
Itil et al 1970(129)	Method: short-term
Itil et al 1971(130)	Method: short-term
Jhee 2003(131)	Method: short-term
Johns 1990(132)	Other: no study, pooled data
JPRN-JapicCTI-050092(133)	Method: short-term
JPRN-JapicCTI-101146(134)	Method: short-term
Kahn et al 2007(135)	Method: short-term
Kane 1993(136)	Method: Short term
Kane et al 2002a(137)	Method: short-term
Kane et al 2003(138)	Method: short-term
Kane et al 2007(139)	Method: short-term
Kane et al 2010a(140)	Method: short-term
Kane et al 2011(141)	Method: randomized
	Population: schizophrenia
	Intervention: flexible
Kane et al 2015(142)	Method: short-term
Kane et al 2015a(143)	Method: short-term
Kapur et al 1998(144)	Method: short-term
Kapur et al 2000(145)	Method: short-term
Kapur et al 2000a(146)	Method: short-term
Karpouzian-Rogers et al 2020(147)	Method: randomized
	Population: schizophrenia
	Intervention: flexibly dosed
Kato et al 2012(148)	Method: randomized
	Population: Schizophrenia
	Intervention: flexible dosages in extension phase
Keck et al 2001(149)	Method: short-term
Keskiner et al 1968(150)	Method: randomized
	Population: schizophrenia
	Intervention: flexible
Khanna et al 1997(151)	Method: randomized
	Population: acutely ill
King et al 1979(152)	Method: short-term
King et al 1998(153)	Method: short-term
Kinon et al 1993(154)	Method: short-term
Kinon et al 2001(155)	Other: only pooled data available
Kinon et al 2004(156)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed dosage
Kinon et al 2008(157)	Method: short-term
Kinon et al 2010(158)	Method: short-term
	Method: Short-term
	Method: short-term
Kileser et al 1996(161)	IVIETNOO: SNOIT-TEIM
KO ET AL 1995(162)	IVIETNOO: SNOIT-TEIM
Kramer et al 2007(163)	IVIETNOO: SNOIT-TEIM
Kryznanovskaya et al 2009(164)	IVIETNOD: SNORT-TERM
Kudo et al 1985(165)	Method: unclear
	Intervention: flexible
Kurland et al 1975(166)	Intervention: flexible Method: short-term

Lan et al 2007(168)	Method: short-term
Landbloom et al 2017(169)	Method: short-term
Lane et al 2001(170)	Method: short-term
Lecrubier et al 1988(171)	Method: short-term
Lecrubier et al 2006(172)	Method: short-term
Lee et al 2002(173)	Method: short-term
Lee et al 2012(174)	Method: short-term
Lehmann et al 1980(175)	Method: short-term
Lesem et al 2001(176)	Method: short-term
Levin et al 1996(177)	Method: short-term
Levinson et al 1992(178)	Method: short-term
Levy et al 1983(179)	Method: short-term
Li et al 2014(180)	Method: short term
Li et al 2014a(181)	Method: short-term
Liebermann et al 2016(182)	Method: short-term
Lindenmayer et al 2009(183)	Method: short-term
Lindenmayer et al 2011(184)	Method: randomized
	Population: chronic schizophrenia
	Intervention Galantamine augmentation
Llaudo et al 2016(185)	Method: short-term
Loebel et al 2013(186)	Method: short-term
Loebel etal 2015(187)	Method: short-term
Louza Neto et al 1988(188)	Method: short-term
Mahal et al 1975(189)	Method: short-term
Mamo et al 2004(190)	Method: short-term
Mamo et al 2007(191)	Method: short-term
Marder et al 1994(192)	Method: randomized
	Participants: schizophrenia
	Intervention: flexibly dosed
Marder et al 2007(193)	Method: short-term
Marinkovic et al 2006(194)	Method: short-term
Martin et al 2019(195)	Method: short-term
Mathur et al 1981(196)	Method: randomized
	Population: chronic schizophrenia
	Intervention: probably flexible doses, unclear
	Outcome: no usable data
Matsumoto et al 2018(197)	Method: randomized
	Population: schizophrenia
	Intervention: flexible dosed
Mauri et al 2006(198)	Method: short-term
Mavroidids et al 1983(199)	Method: short-term
Mavroidis et al 1984(200)	Method: short-term
McClelland et al 1974(201)	Method: randomized
	Population: treatment resistant = not stable
McDonnel et al 2008(202)	Method: short-term
McEvoy et al 1991(203)	Method: short-term
McEvoy et al 1996(204)	Method: short-term
McGorry et al 2011(205)	Method: short-term
Meltzer et al 2011(206)	Method: short-term
Meltzer et al 2012(207)	Method: short-term
Meltzer et al 2014(208)	Method: randomized
<u> </u>	Population: treatment resistant = not stable
Meltzer et al 2015(209)	Method: randomized
	Population: schizophrenics

	Intervention: no monotherapy
Meltzer et al 2015a(210)	Method: short-term
Merlo et al 2002(211)	Method: short-term
Miceli et al 1998(212)	Method: short-term
Mitchell et al 2003(213)	Method: short-term
Mitchell et al 2006(214)	Method: short-term
Modestin et al 1983(215)	Method: short-term
Mosholder et al: Study 0008(216)	Method: short-term
Mosholder et al: Study 0012(216)	Method: short-term
Mosholder et al: Study 0013(216)	Method: short-term
Nair et al 1998(217)	Method: randomized
	Population: treatment refractory =not stable
Nakamura et al 2016(218)	Method: short-term
Nasrallah et al 2010(219)	Method: short-term
Nasrallah et al 2013(220)	Method: short-term
Nasser et al 2016(221)	Method: short-term
NCT 00044044(222)	Method: short-term
NCT 00650611(223)	Method: short-term
NCT00044005(224)	Other: no usable data
NCT00074477(225)	Other: no usable data
NCT00077714	Method: short-term
Published in Meltzer et al 2008(226)	
NCT00078039	Method: short-term
Published in Meltzer et al 2008(226)	
NCT00083668	Method: short-term
Published in Meltzer et al 2008(226)	
NCT00085748(227)	Other: no usable data
NCT00088075(228)	Method: short-term
NCT00210548(229)	Method: short-term
NCT00210717(230)	Method: randomized
	Participants: schizophrenia
	Intervention: flexibly dosed
NCT00232687(231)	Method: short-term
NCT00237939(232)	Method: short-term
NCT00297947(233)	Method: short-term
NCT00485810(234)	Method: short-term
NCT00653406(235)	Method: short-term
NCT00704509(236)	Method: short-term
NCT00711269(237)	Method: short-term
NCT00796081(238)	Method: short-term
NCT00862992(239)	Method: short-term
NCT00892528(240)	Method: short-term
NCT00905307(241)	Method: short-term
NCT01082250(242)	Method: short-term
NCT01142596(243)	Other: no usable data
NCT01377233(244)	Method: short-term
NCT01423916(245)	Method: short-term
NCT01493726(246)	Other: No usable data
NCT01606254(247)	Method: short-term
NCT01625000(248)	Method: short-term
NCT01625897(249)	Method: randomized
	Population: schizophrenia
	Intervention: after 4 first weeks flexibly dosed
NCT01626456(250)	Method: randomized

(Meltzer 2015_Ex)	Participants: not stable
NCT01626859(251)	Method: short-term
NCT01626872(252)	Method: randomized
	Population: schizophrenia, but unclear how
	stable the participants were
	Outcome: no useable data
NCT01942382(253)	Method: short-term
NCT02146547(254)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed doses
NCT02174510(255)	Method: short-term
NCT03751488(256)	Method: short-term
NCT03817502(257)	Method: short-term
NCT03870880(258)	Method: randomized
	Population: probably not stable but (still) acutely
· · · · · · · · · · · · · · · · · · ·	
NCT03872596(259)	Method: short-term
NCT04030143(260)	Method: randomized
	Population: not 80% schizophrenia, also bipolar I
Neborsky et al 1981(261)	Method: short-term
Odejide et al 1982(262)	Method: randomized
0 () () () () () () () () () (Population: acute schizophrenia
Ogasa et al 2013(263)	Method: short-term
Ono et al 2006(264)	Method: short-term
Ono et al 2008(265)	Method: short-term
Oostuizen et al 2004(266)	Method: short-term
Oren et al 2007(267)	Method: short-term
Ortega-Soto et al 1994(268)	Method: short-term
Ota et al 1973(269)	Method: short-term
Palao et al 1994(270)	Method: short-term
Pandina et al $2010(271)$	Method: Short-term
Petrie et al $1997(272)$	Method: short-term
Peuskens et al 2007(273)	Method: Tandomized
	Population. Chronic Schizophrenia
Datkin at al 1085(274)	Method: abort torm
$\frac{1903(274)}{275}$	Method: short term
$\frac{1}{2} = \frac{1}{2} = \frac{1}$	Method: short term
Polkin et al $2003(270)$	Method: short term
$\frac{PO(KIII \text{ et al } 2000(277))}{Potkin \text{ et al } 2012(278)}$	Method: short term
$\frac{1}{2013(270)}$	Method: short term
$\frac{1}{2014(219)}$	Method: short torm
Prion at al 1900(200)	Method: randomized
	Population: chronic schizonbrenia
	Intervention: wrong medication
Puech et al 1998(282)	Method: short-term
$\Omega_{\rm uitkin et al 1975(283)}$	Method: Short term
	Population: treatment resistant = not stable
Ravenstiin et al 2016(284)	Method: randomized
	Population: schizophrenia
	Intervention: single dose design
Rein et al 1996(285)	Method: short-term
Reschke et al 1974(286)	Method: short-term
Rifkin et al 1977(287)	Method: randomized

	Population: chronic schizophrenia
	Intervention: flexible doses
Rifkin et al 1991(288)	Method: short-term
Rodriguez et al 2004(289)	Method: randomized
	Population: chronic schizophrenia or
	schizoaffective disorder
	Intervention: fixed medication doses
	Outcome: no usable data
Roelofs et al 1974(290)	Method: randomized
	Population: chronic schizophrenia
	Intervention: wrong medication
Rossenu et al 2008(291)	Method: short-term
Rui et al 2014(292)	Method: variable duration in double blind
	treatment phase
Ruiz et al 1975(293)	Method [,] short-term
Ruskin et al 1991(294)	Method: randomized
	Population: chronic schizophrenia
	Intervention: no fixed doses individual dosages
Sampath et al 1992(295)	Method: randomized
	Population: chronic schizophrenia
	Intervention: no fixed doses individual dosages
	continued
Santos et al 1989(296)	Method: short-term
Sarin et al 2004(297)	Method: short-term
Schooler et al 2000(298)	Method: short-term
Schooler et al 2000(200)	Method: randomized
Shawver et al 1939(299)	Population: chronic schizonhronia
	Intervention: fixed doses
	Outcome: no relanse provention, no
	modeline. The relapse prevention, no
Sim at al $1080(300)$	Mothod: short-torm
Simpson at al 1067(201)	Method: short torm
Simpson et al $1000(202)$	Method: short-term
	Population: troatmont refractory – not stable
Sinch at al $1000(202)$	Mothod: abort torm
Singh et al 2011(201)	Method, short term
Singh et al 2011(304)	Method: Short-term
Smith et al 1984(305)	Method: Short-term
Smith et al 1987 (306)	Method: short-term
Soria et al 1994(307)	Method, randomized
	Population: chronic schizophrenia
	Intervention: no fixed dose
Stewart et al 2009(308)	Method: randomized
	Population: not 80% schizophrenia
Stone et al 1995(309)	Method: short-term
Sun et al 2018(310)	Method: short-term
Svestka et al 2003(311)	Method: short-term
Switt et al 2002(312)	Method: short-term
Takeuchi et al 2014(313)	Method: randomized
	Population: Schizophrenia
	Intervention: dose reduction by half = no fixed
	dose
I urncliff et al 2012(314)	Method: short-term
Uzun et al 2002(315)	Method: short-term
Van Erp et al 2020(316)	Method: short-term

Van Kammern et al 1996(317)	Method: short-term
Van Putten et al 1986(318)	Method: short-term
Van Putten et al 1990(319, 320)	Method: short-term
Van Putten et al 1991(321)	Method: short-term
Vandecasteele et al 1974(322)	Method: randomized
	Population: schizophrenia
	Intervention: wrong medication, no fixed doses
Vanover et al 2016(323)	Method: short-term
Vanover et al 2016(324)	Method: short-term
Vinar et al 1970(325)	Method: short-term
Walling et al 2018(326)	Method: short-term
Wehnert et al 1999(327)	Method: short-term
Weiden et al 2016(328)	Method: randomized
	Population: chronic schizophrenia
	Intervention: flexible doses
Wessels et al 1991(329)	Method: short-term
Wiles et al 1980(330)	Method: short-term
Winter et al 1984(331)	Method: short-term
Witte et al 2012(332)	Method: short-term
Yamagami et al 1990(333)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed doses
Younis et al 2012(334)	Method: short-term
Zimbroff et al 1997(335)	Method: short-term
Zissis et al 1982(336)	Method: randomized
	Population: schizophrenia
	Intervention: no fixed doses

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In sum there were 340 studies, 26 were included and 314 were excluded. Please note that some publications reported on two studies so that there are only 336 references.

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eAppendix 3. Assessment with the Cochrane Risk of bias tool, version2 : judgements about each bias item for each study for the primary outcome relapse

Study name	Bias arising from the randomisation process	Bias due to deviations from intended interventions	Bias due to missing outcome data	Bias in measurement of the outcome	Bias in selection of the reported results	Overall risk of bias
Arato 2002	Low	Some concerns	Low	Some concerns	Low	Some concerns
Velligan 2002	Low	Low	Low	Low	Low	Low
Beasley 1996b_Extension	Some concerns	Low	Some concerns	Low	Low	Some concerns
Beasley 1997_Extension	Some concerns	Low	High	Low	Low	High
Carpenter 1999	Some concerns	Low	Low	Low	Low	Some concerns
Chen 2010	Low	Some concerns	Low	Some concerns	Low	Some concerns
Cooper 2000b	Low	Some concerns	Low	Some concerns	Low	Some concerns
Dotti 1979	Some concerns	Some concerns	Low	Some concerns	Low	Some concerns
Eklund 1991	Some concerns	Some concerns	Low	Some concerns	Low	Some concerns
Fleischhacker 2014	Low	Low	Low	Low	Low	Low
Hough 2009	Low	Low	High	Low	Low	High

Huttunen 1996	Some concerns	Some concerns	Low	Some concerns	Low	Some concerns
Kane 1979	Some concerns	Some concerns	Low	Some concerns	Low	Some concerns
Kane 1983	Some concerns	Low	High	Low	Low	High
Kane 2002b	Low	Low	Low	Low	Low	Low
Kane 2010c	Low	Low	Low	Low	Low	Low
Kane 2012	Low	Some concerns	Low	Some concerns	Low	Some concerns
Mallikaarjun 2013	Low	Some concerns	High	Some concerns	Low	High
Marder 1984	Some concerns	Low	High	Low	Low	High
McEvoy 2007b_Extension	Some concerns	Low	High	Low	Low	High
Nishikawa 1982	Some concerns	Some concerns	Low	Some concerns	Low	Some concerns
Nishikawa 1984	Some concerns	Some concerns	Some concerns	Some concerns	Low	High
Pigott 2003	Low	Some concerns	Low	Some concerns	Low	Some concerns
Schooler 1993	Some concerns	Some concerns	High	Some concerns	Low	High
Simpson 2006	Low	Low	Low	Low	Low	Low



Explanations for our decisions concerning risk of bias judgements according to the risk of bias tool version 2

For judgement of risk of bias, we followed the concept of the Cochrane Risk of Bias tool 2.(Sterne et al. 2019 and Higgins et al. 2019)

This tool provides a framework for evaluating potential risks of bias in five different domains and provides guidance by signaling questions. However, there are not always clear rules and specific situations found in the analyzed trials may deviate from the ideal case. Thus, judgement is needed to make decisions and these specific judgements and decisions made by the authors of the review are made explicit below.

Domain 1: RANDOMIZATION PROCESS



Algorithm for suggested judgement of risk of bias arising from the randomization process

1.1 Was the allocation sequence random?

In principle, if there was no information about the exact methods (e.g. only stated "randomized"), we stated "not indicated". For trials investigating second-generation-antipsychotic drugs that were sponsored by pharmaceutical companies, we assume that the sequence generation for randomization was appropriate, even when it is only stated "randomized", and we stated "probably yes". The reason is that we contacted many pharmaceutical companies in the past and all reported use of appropriate methods in these modern studies, even when it was not clearly stated in the primary publications.

1.2 Was the allocation sequence concealed until participants were enrolled and assigned to interventions?

Similar to 1.1.

1.3 Did baseline differences between intervention groups suggest a problem with the randomization process?

No specific comments

Domain 2: DEVIATIONS FROM INTENDED INTERVENTIONS



Algorithm for suggested judgement of risk of bias due to deviations from the intended interventions (effect of assignment to intervention)

Part 1:

2.1 Were participants aware of their assigned intervention during the trial?

If only stated "double-blind" without further information about the methods, a judgement is needed. We decided to assume that the method of blinding was appropriate and to state "probably no", as in studies of antipsychotic drugs blinding can be rather easily achieved by encapsulating drugs with identical capsules.

In placebo-controlled trials, following the suggestion of the RoB2-guidance document, (Sterne et al. 2019) we assumed unblinding due to side effects. In head-to-head trials of antipsychotics, we did not make this assumption, because the different antipsychotics still have some similarities (overlapping receptor-binding-profiles). Consequently, differences in side-effects are more difficult to evaluate for patients and personal which makes it more difficult to guess the assigned intervention.

2.2 Were carers and people delivering the interventions aware of participants' assigned intervention during the trial?

Similar to 2.1.

2.3 If Y/PY/NI to 2.1 or 2.2: Were there deviations from the intended intervention that arose because of the experimental context?

This question is only relevant for unblinded studies (open, single-blind or placebo-controlled (unblinded due to side effects) trials).

Typically protocol deviations are not reported in detail, which leads to a judgement of "some concerns".

Although protocol deviations due to the experimental context cannot be excluded, we do not deem that substantial protocol deviations (that potentially affect the outcome, see questions below), happen frequently. Thus, we do not expect important bias from deviations of the outcome and a judgement of "some concerns" seems fair or even too punitive.

2.4. Were these deviations likely to have affected the outcome?

No specific comments

2.5. Were these deviations from intended intervention balanced between groups?

No specific comments

Part 2:

2.6. Was an appropriate analysis used to estimate the effect of assignment to intervention?

We considered completer analyses as inappropriate because from such analyses patients are excluded post-randomization due to toxicity or lack of efficacy.

2.7. Was there potential for a substantial impact (on the results) of the failure to analyse participants in the group to which they were randomized?

According to the guidance, authors need to decide about when exclusion of patients post-randomization could have a substantial impact on the results.

We considered completer analyses at "some concerns" when the total number of patients with premature study discontinuation was at maximum 20% of the number randomized.

We considered completer analyses at "high risk" when more than 20% of the patients randomized discontinued prematurely.

The decision for this threshold was informed by the work of Xia et al. (Xia et al. 2009)

Domain 3: MISSING OUTCOME DATA



Algorithm for suggested judgement of risk of bias due to missing outcome data

3.1 Were data for this outcome available for all, or nearly all, participants randomized?

We used the threshold of 5% (study discontinuation rate at maximum 5% of number of patients randomized) mentioned in the RoB2-guidance-document.

For the outcome "relapse" which we analyzed here, this threshold was applied to the number of participants who discontinued for reasons other than relapse.

3.2 If N/PN/NI to 3.1: Is there evidence that the result was not biased by missing outcome data?

We conducted a sensitivity analysis as suggested by the RoB2-guidance-document using plausible assumptions (i.e. patients who discontinued due to inefficacy were assumed to have a relapse; patients who discontinued for reasons other than inefficacy were assumed to have the same relapse rate as observed in the trial (after counting discontinuation due to inefficacy as relapse)). We considered a result as at low risk of bias, when the result of the sensitivity analyses did not differ more than by a factor of 0.8/1.25 from the observed result (i.e. as an example, it is considered acceptable, when in the sensitivity 23 instead of 20 relapses were assumed to happen in a study group of 100 patients (and the event-rate in the other group did not change)).

3.3 If N/PN to 3.2: Could missingness in the outcome depend on its true value?

If no reasons for study discontinuation are reported, then "probably yes", because in studies of antipsychotics in schizophrenia, discontinuation due to lack of efficacy (potentially related to relapse) are likely.

Also, for many reported reasons, doubts remain whether the reason is related to efficacy.

Moreover, it needs to be noted that in our aggregate data (where events are usually reported from all patients randomized) also patients that discontinued due to reasons unrelated to the outcome can affect the result. This is because patients who discontinued prematurely are not at risk for the event anymore.

Thus, all studies with rates of premature study discontinuation above the threshold mentioned in 3.1 need further evaluation.

3.4 If Y/PY/NI to 3.3: Is it likely that missingness in the outcome depended on its true value?

As recommended in the RoB2-guidance-document, we investigated whether there were differences in the total number of participants with premature study discontinuation (dropouts) and in the number of participants with premature study discontinuation for reasons related to the outcome. Thereby, we judged whether it is likely that missingness depended on the outcome and that missingness influenced the outcome substantially (high risk) or to some extend (some concerns).

For the outcome "relapse" analyzed here, we make use of the sensitivity analysis described in 3.2 which includes information about the total number of dropouts and the number of dropouts for related reasons, i.e. due to inefficacy. When the result of the sensitivity analysis differed less than by a factor of 0.67/1.50, but more then 0.8/1.25, from the observed result, then we judged the result at "some concerns". When the result of the sensitivity analysis differed more than by a factor of 0.67/1.50, we judged the result at "high risk" (i.e. as an example, when in a sensitivity analysis more than 27 instead of 20 relapses were assumed in a study group of 100 participants (and the result of the other group did not change)).

When sensitivity analyses investigating the potential impact of missing data on the results were not possible (because information on study discontinuations was not clear enough), we judged the mechanism of missingness and its potential impact on the result according to the following algorithm:

When the rate of study discontinuation for reasons other than relapse was $\leq 20\%$ (in each arm of a comparison), we judged at some concerns. This threshold was informed by the work of Sacket et al.(Sackett 1998) and Xia et al.(Xia et al. 2009). Otherwise proceed.

When the rate ratio of study discontinuation for reasons other than relapse (between two groups compared in a trial) is <0.5/ >2 (half/double), we judged at high risk. Otherwise proceed.

When the rate of study discontinuation due to related reasons (i.e. due to inefficacy) was $\leq 20\%$ (in each arm of a comparison), we judged at some concerns. Otherwise proceed.

When the rate of study discontinuation due to related reasons (between two groups compared in a trial) is <0.5/>2 (half/double), we judged at high risk of bias, when $\ge 0.5/\le 2$, we judge at some concerns.)

Domain 4: MEASUREMENT OF THE OUTCOME



Algorithm for suggested judgement of risk of bias in measurement of the outcome

4.1. Was the method of measuring the outcome inappropriate?

No specific comments

4.2. Could measurement or ascertainment of the outcome have differed between intervention groups?

No specific comments

4.3. Were outcome assessors aware of the intervention received by study participants?

In head-to-head studies of antipsychotic drugs, when only reported that the study was double-blind, we assumed that blinding was appropriate and stated probably yes (similar to 2.1.).

In open trials or double-blind placebo-controlled trials (with potential unblinding of study personal, see 2.1.) we checked if there were particular methods to blind the outcome assessors. If such particular methods were not explicitly described, we assumed that the outcomes were assessed by study personal and answered "probably yes".

4.4 If Y/PY/NI to 4.3: Could assessment of the outcome have been influenced by knowledge of intervention received?

We considered the outcome "relapse" as potentially influenced by knowledge of the intervention received.

4.5 If Y/PY/NI to 4.4: Is it likely that assessment of the outcome was influenced by knowledge of intervention received?

In general, we considered the influence of knowledge of intervention received as minor, resulting in a judgement of some concerns.

Domain 5: SELECTION OF THE REPORTED RESULTS



Algorithm for suggested judgement of risk of bias in selection of the reported result

5.1. Were the data that produced this result analysed in accordance with a pre-specified analysis plan that as finalized before unblinded outcome data were available for analysis?

Typically, the analysis plan was not available. In this case, we followed the recommendations of the Cochrane handbook(Higgins 2020) and compared the reported results with the reported methods section and with the outcomes that are expected for such trials as informed by other trial.

Is the numerical result being assessed likely to have been selected, on the basis of the results, from...

5.2. ... multiple eligible outcome measurements (e.g. scales, definitions, time points) within the outcome domain?

No specific comments

5.3 ... multiple eligible analyses of the data?

No specific comments

Overall risk of bias:

Overall risk-of-bias judgement	Criteria				
Low risk of bias	The study is judged to be at low risk of bias for all domains for this result.				
Some concerns	The study is judged to raise some concerns in at least one domain for this result, but not to be at high risk of bias for any domain.				
High risk of bias	The study is judged to be at high risk of bias in at least one domain for this result. Or				
	The study is judged to have some concerns for multiple domains in a way that substantially lowers confidence in the result.				

For multi-arm-studies (i.e. with several comparisons), for each domain, we used the average of the different comparisons, for the rating of the study. In case of intermediate averages between two categories (e.g. between "high risk" and "some concerns"), we conservatively used the more severe rating (here "high risk").

We judged a study at overall high risk of bias when at least 1 domain was rated at "high risk" or when 4 or more domains were rated as "some concerns".

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eAppendix 4. Individual drugs and Additional sensitivity/subgroup analyses

Aripiprazole oral and long-acting injectable pooled

Relapse (5 studies, 11 dose arms)



Available studies: Pigott 2003, McEvoy 2007b_extension, Kane 2012, Fleischhacker 2014, Mallikaarjun 2013)





Available studies: Pigott 2003, McEvoy 2007b_extension, Kane 2012, Fleischhacker 2014, Mallikaarjun 2013





Available studies: Kane 2012, Fleischhacker 2014







Available studies: Pigott 2003, Kane 2012, Fleischhacker 2014, Mallikaarjun 2013 © 2021 American Medical Association. All rights reserved. 63 Available studies: Pigott 2003, McEvoy 2007b_extension, Kane 2012, Fleischhacker 2014, Mallikaarjun 2013)

2012, Fleischhacker 2014, Mallikaarjun 2013

Dropouts due to side-effects (5 studies, 11 dose arms)



Aripiprazole oral separately

5

Aripiprazole LAI separately



Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference, LAI = long-acting injectable

Description of the aripiprazole results

First slide, oral and long-acting injectable pooled

Five studies with 11 individual dose arms were available, two on oral aripiprazole (Pigott 2003, McEvoy 2007b_extension), three on aripiprazole long-acting injectable (LAI), all of which used the "maintena" formulation originally developed by Otsuka and BristolMyersSquibb (Kane 2012, Fleischhacker 2014, Mallikaarjun 2013). Long-acting injectable doses were converted to daily doses. The dose response curve was flat at approximately 12.5mg/day for relapse (5 studies) and overall symptoms (5 studies). For rehospitalisation it flattened earlier, but this result was based on only 2 studies. 12.5mg/day is similar to the near-to-maximum dose found for acute treatment (Leucht S, Crippa A, Siafis S, Patel MX, Orsini N, Davis JM. Dose-Response Meta-Analysis of Antipsychotic Drugs for Acute Schizophrenia. Am J Psychiatry. 2020 Apr 1;177(4):342-353). The all-cause discontinuation slope showed an initial increase (i.e. aripiprazole worse) and then a decrease (i.e. aripiprazole better). This is plausible in the sense that most dropouts are due to inefficacy.

Second slide, oral and long-acting injectable separately

The second aripiprazole slide shows that the results for were similar when aripiprazole oral (plateau ~12.5mg/day) and aripiprazole LAI (plateau ~ 270mg monthly \triangleq 10mg/day were analysed separately. Only relapse and overall symptoms were analyzed, because data on other outcomes were scarce.

Fluphenazine long-acting injectable



Relapse (6 studies, 13 dose arms)



Rehospitalisation

Overall symptoms (2 studies, 4 dose arms)



Available studies: Carpenter 1999, Dotti 1979, Kane 1979, Kane 1983, Marder 1984, Schooler 1993

Available studies: Carpenter 1999, Kane 1983, Marder 1984, Schooler 1993

Available studies: Carpenter 1999, Dotti 1979





Dropouts due to side-effects (no study)

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Available studies: Carpenter 1999, Dotti 1979, Kane 1979, Kane 1983, Schooler 1993

Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference

Six studies with 13 individual dose arms were available (Carpenter 1999, Dotti 1979, Kane 1979, Kane 1983, Marder 1984, Schooler 1993), all using long-acting injectable formulations which were converted to biweekly doses. The dose-response curve for relapse does not show a clear plateau. However, the dose-response curves for rehospitalization and all-cause discontinuation were essentially flat at 15mg biweekly. The curve for overall symptoms is difficult to interpret, because only 2 studies reported this outcome and confidence intervals were extremely wide, but it also suggests that 15mg/biweekly is the plateau dose. No study provided data on dropouts due to adverse events/side-effects.



Haloperidol oral and long-acting injectable pooled

Available studies: Beasley 1996_extension, Eklund 1991, Huttunen 1996, Kane 2002, Nishikawa 1982, Nishikawa 1984







Dropouts due to side-effects

(5 studies, 14 arms)



Available studies: Beasley 1996_extension, Huttunen 1996

All cause discontinuation (5 studies, 14 arms)



Available studies: Beasley 1996_extension, Eklund 1991, Huttunen 1996, Kane 2002, Nishikawa 1984



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Available studies: Beasley 1996_extension, Eklund 1991, Kane 2002, Nishikawa 1982, Nishikawa 1984

Relapse - haloperidol oral (3 studies, 8 arms)

Relapse - haloperidol long-acting injectable (3 studies, 8 arms)



Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD standardized mean difference dical Association. All rights reserved. 69

Description of the haloperidol results

First slide, oral and long-acting injectable pooled

Six studies were available, three on long-acting injectable haloperidol (Kane 2002, Huttunen 1996, Eklund 1991) and three on oral haloperidol (Nishikawa 1982, Nishikawa 1984, Beasley 1996). Long-acting injectable doses were converted to daily doses. The dose-response curves for relapse and all-cause discontinuation plateauted at approximately 3mg/day for relapse, but slightly increased thereafter for all-cause discontinuation. For overall efficacy and rehospitalisation only 2 studies were available making the data uninterpretable. Dropouts due to adverse events were higher in the haloperidol groups than in the placebo groups already at low doses and plateatued at approximately 3mg/day. The large confidence intervals needs to be considered.

Second slide, oral and long-acting injectable separately

Separate analyses of oral (plateau ~ 3mg/day) and long-acting injectable formulations (plateau ~75mg monthly \triangleq 2.7mg/day) on the second haloperidol slide showed consistent results. We only analysed relapse, because for the other efficacy outcomes overall symptoms and rehospitalisation only two studies were available

Olanzapine – oral and long-acting injectable pooled

Relapse (3 studies, 12 dose arms)





Rehospitalisation

Overall symptoms (3 studies, 12 dose arms)



Available studies: Kane 2010, Beasley 1996, Beasley 1997

All cause discontinuation (3

studies, 12 dose arms)

Available studies: Kane 2010, Beasley 1996, Beasley 1997

Dropouts due to side-effects (3 studies, 12 dose arms)





Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference

Olanzapine – oral and long-acting injectable studies separately



Oral (2 studies, Beasley 1996, 1997)

Long-acting injectable (1 study, Kane 2010)

Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference
Description of the olanzapine results: Three studies with 12 individual dose arms were available, one on olanzapine long-acting injectable (Kane 2010) and two on oral olanzapine (Beasley 1996 and Beasley 1997). The dose response curve for relapse, rehospitalisation and overall symptpoms and all-cause discontinuation seemed to be flat at around 10mg/day, although they still go slightly down up to the maximum examined dose of 20mg/day, in particular for overall symptoms.

A possible explanation for this phenomenon is that **the two oral studies were "continuation**" studies. This means that acute phase responders (here defined as at least 40% PANSS/BPRS reduction from baseline and outpatient at week 6) were followed up without an additional randomisation. In these studies the efficacy curves flattened at approximately 10mg/day. However, this design has a potential bias in favour of placebo / 1mg/day olanzapine, because it may "corrupt" randomization. Probably only special patients may have shown such a strong response in the acute phase. Thus, the patients in the placebo / 1 mg olanzapine group may differ from those in the other groups.

In contrast, the study **on long-acting injectable olanzapine** by Kane et al. 2010 patients who were stabilized on oral olanzapine were randomized to the various long-acting injectable doses. This study showed no clear efficacy plateau at the highest dose of 300mg biweekly which approximately corresponds to 20mg per day.

The results for oral and long-acting injectable separately are shown on the next page (the results for rehospitalization were similar and can be sent upon request).

Paliperidone long-acting injectable

Relapse (1 study, 3 dose arms) Overall symptoms (1 study, 3 dose arms) All cause discontinuation (1 study, 3 dose arms)



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Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference

Description of the paliperidone long-acting injectable results

One study with three dose arms was available (Hough 2009). The curves for relapse, and all-cause discontinuation flattened at approximately 75mg/four weekly, while the overall symptom curves shows a gradually increasing efficacy up to the maximum examined dose of 100mg/month. The data are uninterpretable for clinical practice, because the maximum licensed dose of 150mg/month was not included in the single study available. Moreover, the enormous uncertainty due to the paucity of data expressed by very wide confidence intervals must be considered.

Quetiapine immediate release

Relapse (2 studies, 5 dose arms)



All cause discontinuation (2 studies, 5 dose arms)



Rehospitalisation (2 studies, 5 dose arms)



Dropouts due to side-effects (2 studies, 5 dose arms)



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Overall symptoms

(0 studies)

Description of the quetiapine results

Two quetiapine <u>immediate</u> release studies (Arvanitis 1993, an unpublished study for which we identified relapse data in a report sent to the FDA; and Chen 2010 which was a first-episode study) were included. They 75mg/day, 300mg/day, 400mg/day (Chen 2010), 600mg/day and placebo. They showed increasing effectiveness in terms of relapse prevention and all-cause discontinuation up to 600mg/day, the maximum dose tested. **It seems that a plateau was not yet reached**. The dose-response curve for rehospitalization remained stable from approximately 300mg/day upward. Data on overall efficacy were not available. Dropouts due to side-effects increased up to a dose of approximately 300mg/day. Data on quetiapine <u>extended</u> release were not available.

Ziprasidone



Footnote for all figures: The marks on the x-axis indicate the available dose arms. OR = odds ratio, SMD = standardized mean difference

One ziprasidone study with four individual dose arms (placebo, 40mg/day, 80mg/day and 160mg/day) was available (Arato 2002). A plateau was reached at approximately 75mg/day for relapse, overall symptoms and all-cause discontinuation. No data on rehospitalisation were available.

Risperidone long-acting injectable

Simpson 2006 compared two doses of risperidone consta, 50mg/14 days was somewhat more efficacious than 25mg/14 days. As dose-response meta-analysis requires at least 3 arms, such an analysis was not possible, but the study was included in the overall analyses across drugs.

Lurasidone

Tandon 2016 compared only a single lurasidone dose with placebo. Therefore, a separate doseresponse meta-analysis was not possible for lurasidone. The study was, however, included in the overall analyses across drugs.

Zotepine

Cooper 2000 compared only a single zotepine dose with placebo. Therefore, a separate doseresponse meta-analysis was not possible for zotepine. The study was, however, included in the overall analyses across drugs. Additional sensitivity and subgroup analyses

Sensitivity analysis of the primary outcome relapse: Eight studies with high overall risk of bias excluded (Beasley

1997_Extension, Hough 2009, Kane 1983, Mallikaarjun 2013, Marder 1984, McEvoy 2007, Nishikawa 1984, Schooler 1993)

Sensitivity analysis of the primary outcome relapse: Two non double-blind studies excluded (Huttunen 1996, Mallikaarjun 2013)



The results of both sensitivity analyses were virtually identical with those of the primary analysis

Subgroup analysis of the outcome relapse: Median split of the mean age of the study population

The post-hoc subgroup analysis dividing the studies in two groups according to the median mean age (38.5 years) is shown below. There was no clear difference between the groups. The older group (red) bulged only slightly earlier, the confidence intervals overlapped broadly and the shape of the curves was similar. One reason may be that the median age in both groups was not very different (\geq median age group: median 41.6 years, <median age group: 35.3 years)



 Mean age: median 38.5 years interquartile range (35.5;42.0)

 Red: mean age ≥ median: 13 studies,36 arms, 3365 particiants

 Median mean age 41.6 IQR (39.5;43.6)

 Studies: Arato 2002, Cooper 2000b, Eklund 1991, Fleischhacker 2014, Hough 2009, Kane 2002b, Kane 2010c, Kane

 2012, Mallikaarjun 2013, Nishikawa 1984, Pigott 2003, Simpson 2006, Tandon 2016

 Blue: mean age < median: 10 studies, 29 arms, 1162 participants</td>

 Median age 35.4 IQR (29.6;35.8)

 Studies: Arvanitis 1993, Beasley 1996b_Extension, Beasley 1997_Extension, Carpenter 1999, Chen 2010, Kane 1979,

 Kane 1983, Marder 1984, Nishikawa 1982, Schooler 1993

Subgroup analysis of the outcome relapse: Median split of the percentage men in the studies

The subgroup analysis dividing the studies in two groups according to the median of percentage male (67%) is shown below. The group with more men (red) bulged earlier (~3mg risperidone equ. per day). The result does not seem to be driven by age which was similar in both subgroups (38.2years versus 39 years). It is unexpected in the sense that one analysis suggested that women respond better than men in the acute phase¹. We would therefore have expected that a lower dose might be sufficient for them. Moreover, two reviews did not find sex to be a predictor of relapse^{2,3}. Moreover, this is a post-hoc analysis, the sample size is small, subgroup analyses are hypothesis generating only, and there maybe hidden moderators explaining this result.



Percentage men: median 67% Interquartile range (60%;80%) Red: Percentage male ≥ median: 12 studies,36 arms, 1150 participants Median percentage male 80% IQR (71%;88%) Age: Median 38.2 years IQR (35.4; 41.6) Studies: Arato 2002, Arvanitis 1993, Beasley 1996b_Extension, Carpenter 1999, Cooper 2000b, Dotti 1979, Kane 1979, Kane 2002b, Mallikaarjun 2013, Marder 1984, Nishikawa 1982, Nishikawa 1984

Blue: Percentage male < median: 12 studies, 31 arms, 3380 participants^{3,13-23}

Median percentage male 60% IQR (58%;66%)

Age: Median 39.0 years IQR (35.8; 41.2)

Studies: Beasley 1997_Extension, Chen 2010, Fleischhacker 2014, Hough 2009, Huttunen 1996, Kane

1983, Kane 2010c, Kane 2012, Pigott 2003, Schooler 1993, Simpson 2006, Tandon 2016

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eAppendix 5. Conversion to absolute rates, relative risks and PANSS units

Conversion of ORs and SMDs

Formulas for the conversion

The conversions were conducted according to the formulas in the Cochrane Handbook:¹

1. ORs to absolute rates and RRs

The conversion of ORs to absolute rates and RRs requires an assumed control risk (ACR). As ACR, we used the metaanalytic point estimate of event rates in the placebo group (for the outcomes relapse, re-hospitalization, dropouts due to any cause, dropouts due to any reason).

Then, we transformed ORs using the equations:

- From OR to RR: $RR = \frac{OR}{(1 ACR * (1 OR))}$
- From OR to absolute rates: Absolute rate = $ACR * \frac{OR}{(1 ACR*(1 OR))}$

The same formulas were used to transform the lower and upper boundaries of the 95% confidence intervals.

2. SMDs to absolute PANSS change scores

The conversion of SMD to absolute PANSS scores require a) a standard deviation of the scale, for which the weighted average of SD of PANSS change scores was used: 16.47 (median 19.47 IQR [11.56-30.46]); and b) the absolute PANSS change score in the placebo group (PANSS at 0 mg), for which we used the meta-analytic point estimate of change in the placebo group.

Then, we transformed SMDs using the equation: PANSS change scores (dose) = PANSS change score (0 mg) + SMD * SD

Estimating event rates and PANSS change scores in placebo groups

1. Relapse

Study	Events	Total		Proportion 95%-CI
Dotti 1979	3	10 -		0.30 [0.07: 0.65]
Schooler 1993	19	49		0.39 [0.25; 0.54]
Tandon 2016	72	141		0.51 [0.43, 0.60]
Cooper 2000b	30	58		0.52 [0.38; 0.65]
Beasley 1996b_Extension	8	13	•	0.62 [0.32; 0.86]
Pigott 2003	100	155		0.65 [0.56; 0.72]
Eklund 1991	16	23		0.70 [0.47; 0.87]
Arato 2002	50	71		0.70 [0.58; 0.81]
Chen 2010	67	89		0.75 [0.65; 0.84]
Kane 2012	102	134		0.76 [0.68; 0.83]
Kane 1979	7	8		- 0.88 [0.47; 1.00]
Nishikawa 1984	12	13	• •	- 0.92 [0.64; 1.00]
Nishikawa 1982	10	10		- 1.00 [0.69; 1.00]
				=
Random effects model		774		0.67 [0.56; 0.76]
Heterogeneity: $I^2 = 84\%$, τ^2	= 0.4364, /	p < 0.01		
			0.2 0.4 0.6 0.8	1

On average, about 67% of the participants in the placebo group relapsed, and ACR was set at 67%.

2. Re-hospitalization



On average, about 18% of the participants in the placebo group had a re-hospitalization, and ACR was set at 18%.

3. Dropouts due to any reason



On average, about 75% of the participants in the placebo group dropped out due to any reason, and ACR was set at 75%.

4. Dropouts due to side effects

Study	Events	Total				Proportion	95%-CI
Tandon 2016	1	141 +				0.01	[0.00; 0.04]
Kane 2012	2	134 +	_			0.01	[0.00; 0.05]
Eklund 1991	0	23 -				0.00	[0.00; 0.15]
Nishikawa 1984	0	13 -				0.00	[0.00; 0.25]
Dotti 1979	0	10 -				0.00	[0.00; 0.31]
Nishikawa 1982	0	10 -				0.00	[0.00; 0.31]
Kane 1979	0	8			_	0.00	[0.00; 0.37]
Cooper 2000b	4	58 -				0.07	[0.02; 0.17]
Chen 2010	7	89 -				0.08	[0.03; 0.16]
Pigott 2003	13	155				0.08	[0.05; 0.14]
Arato 2002	7	71				0.10	[0.04; 0.19]
Beasley 1996b_Extension	3	15				0.20	[0.04; 0.48]
Random effects model		727 🧹	>			0.04	[0.02; 0.08]
Heterogeneity: $I^2 = 67\%$, $\tau^2 =$	= 0.7642,	p = 0.17	1 1	I	I		
		0	0.1 0.2	2 0.3	0.4		

On average, about 4% of the participants in the placebo group dropped out due to side effects, and ACR was set at 4%.

5. Overall symptoms

First, we pooled original PANSS change scores from baseline to endpoint in the placebo group:



Three out of the seven placebo-controlled studies with data for overall symptoms reported original PANSS change scores. On average, there was a 9.62 increase of PANSS scores from baseline to endpoint in placebo groups.

Since the other four studies reported PANSS endpoint scores or BPRS scores, we further evaluated the robustness of the above meta-analysis. First, we transformed other measures to PANSS mean change scores:

- PANSS mean endpoint to PANSS mean change scores (change score = endpoint baseline scores). The same was applied when BPRS endpoint scores were reported.
- BPRS mean change scores to PANSS mean change scores according to an equipercentile linking method²

The weighted average of PANSS change scores (original or transformed) was 8.9 (median 4.5 IQR [1.75-15.6]), which is in line with the meta-analysis of original data. A meta-analysis of original or transformed data was avoided, since further assumptions for the calculation of their standard deviations were required.

References

1. Higgins JP, Thomas J, Chandler J, Cumpston M, Page MJ, Welch VA. Cochrane Handbook for Systematic Reviews of Interventions version 6.0; 2019.

2. Leucht S, Rothe P, Davis JM, Engel RR. Equipercentile linking of the BPRS and the PANSS. *Eur Neuropsychopharmacol* 2013; **23**(8): 956-9.

eAppendix 6. Heterogeneity and small-trial / publication bias

eAppendix 6_Variation-Partition-Coefficients – Relapse (primary outcome)



eAppendix 6_Variation-Partition-Coefficients – All-cause discontinuation



eAppendix 6_Variation-Partition-Coefficients – Dropouts due to adverse events



eAppendix 6_Variation-Partition-Coefficients – Rehospitalization



eAppendix 6_Variation-Partition-Coefficients – Overall symptoms





eAppendix 6_Contour enhanced funnel plot and Egger's test of the primary outcome (relapse)

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eReference

Crippa A, Discacciati A, Bottai M, Spiegelman D, Orsini N. One-stage dose-response meta-analysis for aggregated data. Stat Methods Med Res. 2019;28:1579-1596 Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. BMJ. 1997 Sep 13;315(7109):629-34. Peters JL, Sutton AJ, Jones DR, Abrams KR, Rushton L. Contour-enhanced meta-analysis funnel plots help distinguish publication bias from other causes of asymmetry. J Clin Epidemiol 2008;61:991-996 eAppendix 7. Update of the meta-analysis of Uchida et al. 2011 and additional analyses of doses higher than standard doses (≥5mg risperidoneequivalent per day)

eAppendix 7_A, Relapse - standard dose (= 1 Daily-Defined-Dose or higher) versus low-dose (≥50 % < 1 DDD)

	Low d	ose	Standard	dose		Odds Ratio		Odds	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Rand	lom, 95% Cl	
Arato 1997	29	72	47	135	13.2%	1.26 [0.70, 2.28]		100	-	
Carpenter 1999	14	25	12	25	6.5%	1.38 [0.45, 4.20]				
Dellva - Beasley 1996	1	14	5	31	2.0%	0.40 [0.04, 3.79]		10 11 N	8	
Dellva - Beasley 1997	0	31	5	83	1.2%	0.23 [0.01, 4.22]	20		10	
Hough 2009	24	82	29	170	12.7%	2.01 [1.08, 3.74]				
Inderbitzin 1994	5	23	4	20	4.2%	1.11 [0.25, 4.87]		19	-	
Kane 2002	11	30	13	60	7.9%	2.09 [0.80, 5.49]		36		
Mallikaarjun 2013	3	27	0	14	1.2%	4.14 [0.20, 86.05]		12		205
McEvoy 2007 extension	12	88	25	88	10.3%	0.40 [0.19, 0.86]			2	
Nishikawa I	4	12	2	12	2.7%	2.50 [0.36, 17.32]		400	0 5 04	
Nishikawa II	10	13	5	11	3.2%	4.00 [0.69, 23.09]		500		
Schooler 1997	16	52	8	55	8.0%	2.61 [1.01, 6.77]				
Simpson 2006	42	162	32	161	14.5%	1.41 [0.84, 2.38]		125		
Velligan 1993	64	88	48	87	12.5%	2.17 [1.15, 4.07]			20-19-02-02	
Total (95% CI)		719		952	100.0%	1.46 [1.04, 2.04]			•	
Total events	235		235							
Heterogeneity: Tau ² = 0.1	3; Chi ² = 3	20.87. 0	#f = 13 (P =	0.08); 1*	² = 38%		L		<u> </u>	
Test for overall effect: Z =	2.20 (P =	0.03)	1	0			0.01	U.1 Favours low dose	1 10 Favours standard do	100 se

Uchida et al. 2011¹ had reported in a pairwise meta-analysis that standard doses (= 1 Daily-Defined-Dose) were just not statistically significantly more effective

than low doses (\geq 50 % < 1 DDD) for relapse prevention with antipsychotics in schizophrenia (p=0.05). We updated their Figure 4a with new studies found by us. We also obtained additional data on studies which had already been included by them (Velligan 1993, Carpenter 1999, Beasley 1996, Beasley 1997, Hough 2009, Kane 2010, Mallikaarjun 2013, McEvoy 2007 extension). After this update standard doses outperformed low doses more clearly (p=0.03). **Comment**

1. Minor differences in numbers between our and their analysis stem from the fact that Uchida et al. 2011¹ a) had several times dropout due to inefficacy/relapse for their analysis while we always used relapse if available; b) they did not always use the strict intention-to-treat (once-randomized – analyzed) population for the denominator. Moreover, in the Schooler 1997 study we used only the arms with 'supportive' family treatment, while they also used the arms with 'intensive' family treatment. Our decision was made *a priori*; and the rationale was that we were interested in the "pure" dose-effect which could have been confounded by additional intensive family treatment (or at least more than by just supportive family treatment).

2. It should be noted that the only three studies which showed an effect in favour of the low dose groups were the only continuation studies (Beasley 1996, Beasley 1997, McEvoy 2007). In this design the responders in the acute phase are followed up. Removing these studies, heterogeneity drops to an l^2 of 0% and the p-value for efficacy is < 0.0001).

3. Most importantly, it should be noted that p-values must not be overemphasized, and there is a strong movement of statisticians against them. The direction of effect in Uchida et al. 2011¹ was in favour of standard doses as well, just the p-value was somewhat higher (also see American Statistical Association 2016²)

References

- 1. Uchida H, Suzuki T, Takeuchi H, Arenovich T, Mamo DC. Low dose vs standard dose of antipsychotics for relapse prevention in schizophrenia: metaanalysis. Schizophr Bull. 2011;37(4):788-799.
- 2. Americian Statistical Association. Statement on statistical significance and p-values. 2016; https://www.amstat.org/asa/files/pdfs/p-valuestatement.pdf

As the slope in Figure 1 of the manuscript – dose-response of the primary outcome relapse – is still slightly increasing above 5mg/day risperidone equivalent, we conducted two post-hoc analyses following reviewer comments:

- 1. An analysis using linear splines as in our dose-response analysis of acute phase studies (Leucht et al. 2020) to find out above which dose the slope of the curve does not longer increase (Figure 2 below)
- 2. A simple pairwise meta-analysis in which we compared doses between 3-7mg/day risperidone equivalent with higher doses (Figure 3 below)

Neither one suggested important efficacy gains at very high doses

There was no significant difference (p=0.48). It should noted that too few 5mg doses were available, therefore we had to use a range. We tested with linear splines up to which dose the dose-response curve still showed a significantly increasing slope (post-hoc, p<0.1)

eAppendix 7, B_ Relapse - above which dose does the slope of the dose-response curve no longer significantly increase?



The red line represents the linear spline at 3.5 mg/day risperidone equivalents, above which dose the slope did not show a significant decrease (p.0.1). The marks on the x-axis indicate for which doses study-arms were available and how many.

eAppendix 7_C Pairwise meta-analysis of OR for relapse of 3-7mg/day risperidone equivalent versus more than 7mg/day

1



We conducted a pairwise meta-analysis in which we compared doses between 3-7mg/day risperidone equivalent with higher doses. Therewas no significant difference (p=0.48). It should be noted that too few arms with exactly 5mg were available, therefore we had to use a range.

Dose, mg	Dopamine receptor
	occupancy, % (95%CI) ^a
Placebo	0 (0 to 0)
0.5	31 (15 to 39)
1.0	45 (31 to 56)
1.5	54 (40 to 64)
2.0	60 (47 to 69)
2.5	64 (54 to 72)
3.0	68 (59 to 75)
3.5	71 (62 to 77)
4.0	73 (65 to 78)
4.5	74 (67 to 80)
5.0	76 (69 to 82)
5.5	78 (71 to 83)
6.0	79 (72 to 84)
6.5	79 (73 to 84)
7.0	80 (73 to 85)
7.5	81 (74 to 86)
8.0	82 (74 to 86)
8.5	82 (75 to 87)
9.0	83 (75 to 88)
9.5	83 (76 to 88)
10.0	84 (76 to 89)
15.0	86 (77 to 92)

eTable 3. Average Dopamine Receptor Occupancies for Risperidone Doses

^aData on dopamine occupancy per risperidone dose were taken from the meta-analysis by Lako et al for use in the discussion section. The median and interquartile ranges for study durations for the various outcomes were as follows: relapse, 48.0 (interquartile range [IQR], 33.0-52.0) months; rehospitalization, 52.00 (IQR, 46.00-52.00) months; efficacy, 46.00 (IQR, 26.00-52.00) months; all-cause discontinuation, 48.00 (IQR, 28.00-52.00) months; and dropout due to adverse events, 46.00 (IQR, 38.00-52.00).

eReference

Lako IM, van den Heuvel ER, Knegtering H, Bruggeman R, Taxis K. Estimating dopamine D2 receptor occupancy for doses of 8 antipsychotics: a meta-analysis. J Clin Psychopharmacol. 2013;33(5):675-681.