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Pregabalin add-on for drug-resistant focal epilepsy (Review)

Panebianco M, Bresnahan R, Marson AG

Panebianco M, Bresnahan R, Marson AG. Pregabalin add-on for drug-resistant focal epilepsy. *Cochrane Database of Systematic Reviews* 2022, Issue 3. Art. No.: CD005612. DOI: 10.1002/14651858.CD005612.pub5.

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[Intervention Review]

Pregabalin add-on for drug-resistant focal epilepsy

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Editorial group: Cochrane Epilepsy Group. **Publication status and date:** Edited (no change to conclusions), published in Issue 4, 2022.

Citation: Panebianco M, Bresnahan R, Marson AG.Pregabalin add-on for drug-resistant focal epilepsy. *Cochrane Database of Systematic Reviews* 2022, Issue 3. Art. No.: CD005612. DOI: 10.1002/14651858.CD005612.pub5.

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ABSTRACT

Background

This is an updated version of the Cochrane Review last published in Issue 7, 2019; it includes two additional studies.

Epilepsy is a common neurological disease that affects approximately 1% of the UK population. Approximately one-third of these people continue to have seizures despite drug treatment. Pregabalin is one of the newer antiepileptic drugs that has been developed to improve outcomes. In this review we summarised the current evidence regarding pregabalin when used as an add-on treatment for drug-resistant focal epilepsy.

Objectives

To assess the efficacy and tolerability of pregabalin when used as an add-on treatment for drug-resistant focal epilepsy.

Search methods

For the latest update we searched the following databases on 16 November 2020: Cochrane Register of Studies (CRS Web), and MEDLINE (Ovid, 1946 to 16 November 2020). CRS Web includes randomised or quasi-randomised, controlled trials from PubMed, Embase, ClinicalTrials.gov, the World Health Organisation International Clinical Trials Registry Platform (ICTRP), the Cochrane Central Register of Controlled Trials (CENTRAL), and the Specialised Registers of Cochrane Review Groups, including Epilepsy.

We imposed no language restrictions. We contacted the manufacturers of pregabalin and authors in the field to identify any relevant unpublished studies.

Selection criteria

We included randomised controlled trials comparing pregabalin with placebo or an alternative antiepileptic drug as an add-on for people of any age with drug-resistant focal epilepsy. Double-blind and single-blind trials were eligible for inclusion. The primary outcome was 50% or greater reduction in seizure frequency; secondary outcomes were seizure freedom, treatment withdrawal for any reason, treatment withdrawal due to adverse effects, and proportion of individuals experiencing adverse effects.

Data collection and analysis

Two review authors independently selected trials for inclusion and extracted the relevant data. Primary analyses were intention-to-treat (ITT). We presented summary risk ratios (RRs) and odds ratios (ORs) with 95% confidence intervals (CIs). We evaluated dose response in regression models. We carried out a risk of bias assessment for each included study using the Cochrane risk of bias tool and assessed the overall certainty of evidence using the GRADE approach.



Main results

We included 11 randomised controlled trials (3949 participants). Nine trials compared pregabalin to placebo. For the primary outcome, participants randomised to pregabalin were significantly more likely to attain a 50% or greater reduction in seizure frequency compared to placebo (RR 1.95, 95% CI 1.40 to 2.72, 9 trials, 2663 participants, low-certainty evidence). The odds of response doubled with an increase in dose from 300 mg/day to 600 mg/day (OR 1.99, 95% CI 1.74 to 2.28), indicating a dose-response relationship. Pregabalin was significantly associated with seizure freedom (RR 3.94, 95% CI 1.50 to 10.37, 4 trials, 1125 participants, moderate-certainty evidence). Participants were significantly more likely to withdraw from pregabalin treatment than placebo for any reason (RR 1.33, 95% CI 1.10 to 1.60; 9 trials, 2663 participants; moderate-certainty evidence) and for adverse effects (RR 2.60, 95% CI 1.86 to 3.64; 9 trials, 2663 participants; moderate-certainty evidence).

Three trials compared pregabalin to three active-control drugs: lamotrigine, levetiracetam and gabapentin. Participants allocated to pregabalin were significantly more likely to achieve a 50% or greater reduction in seizure frequency than those allocated to lamotrigine (RR 1.47, 95% CI 1.03 to 2.12; 1 trial, 293 participants) but not those allocated to levetiracetam (RR 0.94, 95% CI 0.80 to 1.11; 1 trial, 509 participants) or gabapentin (RR 0.96, 95% CI 0.82 to 1.12; 1 trial, 484 participants). We found no significant differences between pregabalin and lamotrigine for seizure freedom (RR 1.39, 95% CI 0.40 to 4.83). However, significantly fewer participants achieved seizure freedom with add-on pregabalin compared to levetiracetam (RR 0.50, 95% CI 0.30 to 0.85). No data were reported for this outcome for pregabalin versus gabapentin. We detected no significant differences in treatment withdrawal rate for any reason or due to adverse effects, specifically, during either pooled analysis or subgroup analysis. Ataxia, dizziness, somnolence, weight gain, headache and fatigue were significantly associated with pregabalin than in active control.

We rated the overall risk of bias in the included studies as low or unclear due to the possibility of publication bias and lack of methodological details provided. We assessed all the studies to be at a high risk of funding bias as they were all sponsored by Pfizer. We rated the certainty of the evidence as very low to moderate using the GRADE approach.

Authors' conclusions

For people with drug-resistant focal epilepsy, pregabalin when used as an add-on treatment was significantly more effective than placebo at producing a 50% or greater seizure reduction and seizure freedom. Results demonstrated efficacy for doses from 150 mg/day to 600 mg/day, with increasing effectiveness at 600 mg doses, although there were issues with tolerability at higher doses. However, the trials included in this review were of short duration, and longer-term trials are needed to inform clinical decision-making. This review focused on the use of pregabalin in drug-resistant focal epilepsy, and the results cannot be generalised to add-on treatment for generalised epilepsies. Likewise, no inference can be made about the effects of pregabalin when used as monotherapy.

PLAIN LANGUAGE SUMMARY

Pregabalin add-on for drug-resistant focal epilepsy

This is an update of a review previously published in 2019.

Background

Epilepsy is a common chronic neurological disease that affects approximately 1% of people in the UK. Approximately 1 in 400 people with epilepsy have seizures that continue despite antiepileptic drug treatment (drug-resistant epilepsy). A number of new antiepileptic drugs have been developed to treat epilepsy, of which pregabalin is one. Use of pregabalin in combination with other antiepileptic drugs can reduce the frequency of seizures, but has some adverse effects.

Aim of review

This review aimed to assess the effectiveness and tolerability of pregabalin when used as an add-on antiepileptic drug in treatmentresistant focal epilepsy.

Study characteristics

This review examined data from eleven trials, including a total of 3949 participants. Study participants were assigned using a random method to take pregabalin, placebo, or another antiepileptic drug in addition to their usual antiepileptic drugs.

Key results

Participants taking pregabalin were more than twice as likely to have their seizure frequency reduced by 50% or more during a 12week treatment period compared to those taking placebo, and were nearly four times more likely to be completely free of seizures. Pregabalin was shown to be effective across a range of doses (150 mg to 600 mg), with increasing effectiveness at higher doses. There was also an increased likelihood of treatment withdrawal with pregabalin. Side effects associated with pregabalin included ataxia, dizziness, fatigue, somnolence, headache, nausea and weight gain. When we compared pregabalin to three other antiepileptic drugs (lamotrigine,



levetiracetam, and gabapentin), participants taking pregabalin were more likely to achieve a 50% reduction in seizure frequency than those taking lamotrigine. We found no significant differences between pregabalin and levetiracetam or gabapentin as add-on drugs.

Certainty of the evidence

We rated all included studies as being at low or unclear risk of bias due to missing information about the methods used to conduct the trial and a suspicion of publication bias. Publication bias can occur when studies that report non-significant findings are not published. We suspected publication bias because the majority of included studies showed significant findings and were sponsored by the same drug company. We assessed the certainty of the evidence for the primary outcome of reduction in seizure frequency as low, meaning that we cannot be certain that the finding reported is accurate. However, we rated the certainty of the evidence for the outcomes seizure freedom and treatment withdrawal as moderate, so we can be fairly confident that these results are accurate. There were no data regarding the longer-term effectiveness of pregabalin, and the use of pregabalin in drug-resistant generalised epilepsy, which should be investigated in future studies.

The evidence is current to 16 November 2020.

SUMMARY OF FINDINGS

Summary of findings 1. Pregabalin compared to placebo for drug-resistant focal epilepsy

Pregabalin compared to placebo for drug-resistant focal epilepsy

Patient or population: drug-resistant focal epilepsy

Setting: outpatient setting

Intervention: pregabalin

Comparison: placebo

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Outcomes	Anticipated absolute effects* (95% CI)		•		•		Relative effect (95% CI)	№ of partici- pants (studies)	Certainty of the evidence (GRADE)	Comments
	Risk with placebo	Risk with pre- gabalin		(studies)						
50% or greater reduction in seizure frequency - ITT	Study populati	on	RR 1.95 (1.40 to 2.72)	2663 (9 RCTs)	⊕⊕⊝⊝ Iow ^a ,b,c,e,f	Pregabalin may increase the proportion of people achieving 50% or greater reduction				
analysis	199 per 1000	388 per 1000	(1.10 (0 2.12)	(3 (13)	(OW ")") -) -).	in seizure frequency according to ITT analy- sis, but we are uncertain.				
Follow-up: range 12 to 17 weeks	p: range 12 to 17 (279 to 543)					sis, but we are uncertain.				
50% or greater reduction in seizure frequency - best-			RR 2.91 (1.92 to 4.42)	2663 (9 RCTs)	⊕⊝⊝⊝ very low a,b,c,e	Pregabalin may increase the proportion of people achieving 50% or greater reduction				
case analysis	199 per 1000 579 per 1000		- (1.32 (0 4.42)	(5 (C13)	very low ^{a,D,C,e}	in seizure frequency according to best-case				
Follow-up: range 12 to 17 weeks		(382 to 880)				analysis, but we are very uncertain.				
50% or greater reduc- tion in seizure frequency -	Study population		RR 1.10 - (0.92 to	2663 (9 RCTs)	⊕⊝⊝⊝ very low ^{a,b,c,e}	Pregabalin might have no effect on the pro- portion of people achieving 50% or greater				
worst-case analysis	344 per 1000	378 per 1000	1.31)	(5 KCTS)	very low ^{a,D,c,e}	reduction in seizure frequency according to				
Follow-up: range 12 to 17 weeks	up: range 12 to 17 (316 to 451)		,			worst-case analysis, however, we are very uncertain.				
Seizure freedom			RR 3.94	1125		Pregabalin likely increases the number of people achieving seizure freedom.				
Follow-up: 12 weeks	11 per 1000	43 per 1000 (16 to 114)	- (1.50 to 10.37)	(4 RCTs)	moderate ^{a,d,e}	איז				
Treatment withdrawal for any	Study populati	on	RR 1.33 (1.10 to	2663 (9 RCTs)	⊕⊕⊕⊝ moderate ^a	Pregabalin likely slightly increases the num- ber of people who withdraw from treatment				



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reason Follow-up: range 12 to 17 weeks	145 per 1000	193 per 1000 (159 to 232)	1.60)			for any reason. However, this effect may or may not be important.		
Treatment withdrawal due to	Study populati	on	RR 2.60	2663 (9 RCTs)	⊕⊕⊕⊝ modorate∂.d.e	Pregabalin likely increases the number of people withdrawing from treatment due to		
adverse effects Follow-up: range 12 to 17 weeks	44 per 1000 114 per 1000 (82 to 160) 3.64) w-up: range 12 to 17 44 per 1000 3.64)	moderates	adverse effects.					
Adverse effect: ataxia	Study populati	on	RR 3.90	1868		Pregabalin probably slightly increases the		
Follow-up: range 12 to 17 weeks			proportion of participants who experienc ataxia.					
Adverse effect: dizziness	Study populati	ulation RR 3.15 (2.23 to	2193 (7 RCTs)	⊕⊕⊝⊝ low ^{a,e}	Pregabalin probably slightly increases the proportion of participants who experience			
Follow-up: range 12 to 17 weeks	85 per 1000	268 per 1000 (189 to 377)	4.44)	(11(C13)	low ^{e,e}	dizziness.		
Adverse effect: fatigue	erse effect: fatigue Study population	RR 1.35 (0.94 to	2448 (8 RCTs)	⊕⊕⊝⊝ lowa,e	Pregabalin probably slightly increases the proportion of participants who experience			
Follow-up: range 12 to 17 weeks	84 per 1000	113 per 1000 (79 to 162)	1.93)	(8 (CTS)	lowaye	fatigue.		
Adverse effect: headache	Study populati	on	RR 0.65	1850	⊕⊝⊝⊝ very low ^{a,b,c}	RR < 1 indicates that nausea is less likely in the pregabalin group.		
Follow-up: range 12 to 17 weeks	.2 to 17 136 per 1000 88 per 1000 (61 to 128) 0.94) 0.94)	very low ^{a,o,c}	the pregabatin group.					
Adverse effect: nausea	Study populati	on	RR 1.20 (0.56 to	1267 (4 RCTs)	⊕⊕⊝⊝ Iow ^{a,e}	Pregabalin probably slightly increases the proportion of participants who experience		
Follow-up: range 12 to 17 weeks	40 per 1000	48 per 1000 (22 to 103)	2.58)	(4 (CTS)	lowaye	proportion of participants who experience nausea.		
Adverse effect: somno- lence	Study populati	on	RR 2.05 (1.49 to	2663 (9 RCTs)	⊕⊕⊝⊝ Iow ^{a,e}	Pregabalin probably slightly increases the proportion of participants who experience		
Follow-up: range 12 to 17 weeks	87 per 1000	178 per 1000 (130 to 244)	2.81)	(,		somnolence.		

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Iverse effect: weight inStudy populationRR 4.35 (2.34 to) 2488 (8 RCTs) $\oplus \oplus \odot \odot$ low ^{a,e}			Pregabalin probably slightly increases the proportion of participants who experience			
Follow-up: range 12 to 17 weeks	22 per 1000	96 per 1000 (51 to 178)	8.11)	(0 (13)		weight gain.
*The risk in the interventic its 95% CI). CI: confidence interval; ITT:					ne comparison grou	p and the relative effect of the intervention (and
GRADE Working Group grad High certainty: We are very Moderate certainty: We are substantially different. Low certainty: Our confider Very low certainty: We have	confident that the t moderately confidence in the effect esti	ent in the effect esti mate is limited: the	mate: the true effe true effect may be	ct is likely to be cl substantially diffe	erent from the estim	
ot provide information on m Downgraded twice for incon Downgraded once for publica	sistency: significant	heterogeneity (P < 0				
Downgraded once for impred Jpgraded once for large effe Jpgraded once for dose resp	tision: number of ev ct: risk ratio was gre onse: dose-response	vents reported (< 400 eater than 2.00. e relationship was c	onfirmed by regres	sion model.		
Downgraded once for impred Upgraded once for large effe Jpgraded once for dose resp ummary of findings 2.	tision: number of ev ct: risk ratio was gre onse: dose-response Pregabalin compa tive comparator fo	vents reported (< 400 eater than 2.00. e relationship was c ared to active cor or drug-resistant fo	onfirmed by regres	sion model.		
Downgraded once for impred Upgraded once for large effe Upgraded once for dose resp Summary of findings 2. I Pregabalin compared to ac Patient or population: drug Setting: outpatient setting Intervention: pregabalin Comparison: active compared	tision: number of ev ct: risk ratio was gre onse: dose-response Pregabalin compa tive comparator fo g-resistant focal epil	vents reported (< 400 eater than 2.00. e relationship was c ared to active cor or drug-resistant fo	onfirmed by regres	sion model.		
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Downgraded once for impred Upgraded once for large effe Upgraded once for dose resp Summary of findings 2. I Pregabalin compared to ac Patient or population: drug Setting: outpatient setting Intervention: pregabalin Comparison: active compared	tision: number of ev ct: risk ratio was gre onse: dose-response Pregabalin compa- tive comparator for g-resistant focal epil rator (gabapentin, la Anticipated abso	vents reported (< 400 eater than 2.00. e relationship was c ared to active cor or drug-resistant fo lepsy amotrigine and leve	onfirmed by regres nparators for dr cal epilepsy tiracetam) Relative effect	ug-resistant for	certainty of	Comments

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Follow-up: range 16 to 21 weeks	491 per 1000	505 per 1000 (417 to 613)				ple achieving a 50% or greater reduction in seizure frequency compared to other active comparators. We are, however, uncertain about this finding.
50% or greater reduc- tion in seizure frequency	Study populati	on	RR 1.60 (1.17 to 2.19)	1286 (3 RCTs)	⊕⊝⊝⊝ very low ^{a,c}	According to best-case analysis, pregabalin may increase the proportion of participants
- best-case analysis	491 per 1000	785 per 1000 (574 to 1000)		. ,	,	achieving a 50% or greater reduction in seizure frequency. However, we are very un-
Follow-up: range 16 to 21 weeks		(374 (0 1000)				certain about this finding.
50% or greater reduc- tion in seizure frequency	Study populati	Study population		1286 (3 RCTs)	⊕⊕⊕⊝ moderate ^a	According to worst-case analysis, prega- balin may decrease the proportion of partici-
- worst-case analysis	732 per 1000	490 per 1000	– (0.62 to 0.74)	(5 (C13)	moderate	pants achieving a 50% or greater reduction in
Follow-up: range 16 to 21 weeks		(454 to 542)				seizure frequency. We are moderately certair about this finding.
Seizure freedom	Study populati	on	RR 0.59 (0.37 to 0.95)			Pregabalin may reduce the number of people achieving seizure freedom, but we are very
Follow-up: range 16 to 21 weeks	106 per 1000	1000 63 per 1000 (39 to 101)	- (0.37 10 0.93)	(2 KC13)	very low ^{a,d}	uncertain.
Treatment withdrawal for any reason	Study populati	on	RR 0.93 (0.76 to 1.13)	1286 (3 RCTs)	⊕⊕⊙⊝ low ^{a,e}	Pregabalin does not appear to affect the number of participants withdrawing from
Follow-up: range 16 to 21 weeks	241 per 1000	224 per 1000 (183 to 273)	(00 to 1.1.0)	(0.1010)		treatment for any reason. However, we are uncertain.
Treatment withdrawal for adverse events	Study populati	on	RR 1.04 (0.73 to 1.48)	1286 (3 RCTs)	⊕⊕⊝⊝ Iowa,e	Pregabalin does not appear to affect the number of participants withdrawing from
Follow-up: range 16 to 21 weeks	85 per 1000	88 per 1000 (62 to 125)	- (0.15 10 1.10)	(3 (613)	lowaye	treatment for adverse effects, but we are un- certain.
Adverse effect: ataxia	Study populati	on	RR 1.72 (0.54 to 5.55)	293 (1 RCT)		Pregabalin probably slightly increases the proportion of participants who experience
Follow-up: range 16 to 21 weeks	50 per 1000	86 per 1000	- (0.57 (0.5.55)	(1 101)	low ^{a,e}	ataxia.
		(27 to 277)				
Adverse effect: dizziness	Study populati	on	RR 1.64 (0.85 to 3.16)	1286		Pregabalin probably slightly increases the proportion of participants who experience
Follow-up: range 16 to 21 weeks	111 per 1000	191 per 1000	- (0.05 (0 5.10)	(3 RCTs)	low ^{a,e}	dizziness.

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		(94 to 351)				
Adverse effect: fatigue			RR 1.72 (0.77 to 3.83)	293 (1 RCT)		Pregabalin probably slightly increases the proportion of participants who experience fa-
Follow-up: range 16 to 21 weeks	99 per 1000	170 per 1000	- (0.77 to 5.85)	(IRCI)	low ^{a,e}	tigue.
		(76 to 379)				
Adverse effect: headache	headache Study population		RR 0.83 (0.41 to 1.65)	1286 (3 RCTs)	0000 d	RR < 1 indicates that headache is less likely in the pregabalin group
Follow-up: range 16 to 21 weeks	119 per 1000	99 per 1000	- (0.41 (0 1.03)	(3 KCTS)	very low ^{a,d}	the pregabatili group
		(49 to 196)				
Adverse effect: nausea	Study population		RR 0.20	509 (1 RCT)	000	RR < 1 indicates that nausea is less likely in the pregabalin group.
Follow-up: range 16 to 21 weeks	59 per 1000	12 per 1000	- (0.04 to 1.01)	(IRCI)	very low ^{a,d}	the pregabalin group.
weeks		(2 to 59)				
Adverse effect: somno- lence	Study populati	Study population RR 1.16 (0.88 to 1.53)		1286 (3 RCTs)	⊕⊕⊝⊝ low ^{a,e}	Pregabalin probably slightly increases the proportion of participants who experience
Follow-up: range 16 to 21	191 per 1000	221 per 1000	- (0.00 to 1.00)	(3 ((3))	lowa	somnolence.
weeks		(168 to 292)				
Adverse effect: weight gain	Study populati	ion	RR 2.87 - (0.94 to 8.75)	1286 (3 RCTs)	⊕⊕⊝⊝ Iow ^{a,e}	Pregabalin probably slightly increases the proportion of participants who experience
Follow-up: range 16 to 21	33 per 1000	95 per 1000	- (0.3+ (0 0.13)	(3 (613)	low ^{a,e}	weight gain.
weeks		(31 to 289)				

*The risk in the intervention group (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; ITT: intention-to-treat; RCT: randomised controlled trial; RR: Risk ratio

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

^aDowngraded once for risk of bias: one study did not confirm the method of randomisation; all studies failed to specify method of allocation concealment; one study did not provide information on method of blinding.

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brary

^bDowngraded once for inconsistency: significant heterogeneity (P < 0.10) was detected within the data set. ^cDowngraded twice for inconsistency: very significant heterogeneity (P < 0.05) was detected within the data set. ^dDowngraded twice for imprecision: very low number of events (< 100) which did not suffice the optimal information size. ^eDowngraded once for imprecision: very low number of events (< 400) which did not suffice the optimal information size.

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9



BACKGROUND

This is an update of a Cochrane Review previously published in 2019 (Panebianco 2019); it includes two additional studies.

Description of the condition

Epilepsy is a common neurological chronic condition characterised by recurrent, unprovoked seizures, that affects approximately 1% of the UK population (Hauser 1990). A single antiepileptic drug (AED) (monotherapy) can induce remission for the majority of those diagnosed. However, up to 30% of people with epilepsy fail to respond to monotherapy (Cockerell 1995). People who have failed to respond to a minimum of two AEDs given as monotherapy are considered to be 'drug-resistant'. The majority of those who are drug-resistant have focal onset (also called focal- or localisationrelated) seizures. During focal-onset seizures, abnormal electrical activity initiates in one part of the brain, and during the course of the seizure the abnormal electrical activity either remains localised or spreads to other parts of the brain (Ramaratnam 2016). For individuals with drug-resistant focal epilepsy, recurrent seizures can reduce quality of life, and may also lead to injuries, social isolation, and depression (Villeneuve 2004). Individuals with this neurological condition pose a significant therapeutic problem, which has led to the development of new AEDs as well as exploration of non-pharmacological treatment options, such as vagal nerve stimulation and epilepsy surgery (Panebianco 2015; West 2015). Over the past two decades, the introduction of several new antiepileptic drugs that are often better tolerated and more manageable than older AEDs has improved the ability to treat individuals with epilepsy. Recent studies have reported that 12% to 17% of treatment-resistant individuals become seizure-free with the addition of a previously untried, in most cases new-generation, antiepileptic drug (Granata 2009).

Description of the intervention

Since the 1990s, numerous new AEDs have become available that aim to provide more potent and better-tolerated treatments for epilepsy. Pregabalin is one of these new compounds with antiepileptic, analgesic, and anxiolytic (anxiety-reducing) properties. Pregabalin has favourable pharmacokinetics: it is not protein bound, is 90% bioavailable, and reaches peak plasma concentrations within 1.5 hours of administration of an oral dose. With repeated doses, a steady state is achieved within 24 to 48 hours. Furthermore, 90% of the drug is eliminated, unmetabolised, by the kidneys, and it has no known drug interactions (Brodie 2005). Pregabalin was launched in the UK market in 2004 as an add-on AED for focal-onset seizures, as well as a treatment for neuropathic pain, and as an anxiolytic in 2006.

How the intervention might work

Pregabalin is an alpha-2-delta ligand that is approved in multiple countries worldwide as an add-on therapy for focal onset seizures in adult and paediatric populations. Pregabalin is structurally related to both the neurotransmitter γ -aminobutyric acid (GABA), and the older antiepileptic drug, gabapentin. Similar to gabapentin, the primary mechanism underlying the pharmacological action of pregabalin does not appear to involve the GABA system. In particular, pregabalin does not bind to GABA-A, GABA-B, or benzodiazepine receptors. Pregabalin is neither metabolically converted to GABA or to a GABA agonist, nor does it have any effect on the uptake or degradation of GABA. In fact, the

primary mode of action of pregabalin is via the inhibition of depolarisation-induced calcium influx at P-, Q-, and N-type voltagegated calcium channels, located at the nerve terminals. At the molecular level, this action is achieved by pregabalin binding to the α -2- δ subunit of voltage-gated calcium channels (Ben-Menachem 2004). As a consequence of the reduced calcium influx, less excitatory neurotransmitter, such as glutamate, is released from the presynaptic nerve terminals.This action is thought to mediate its antiepileptic, anxiolytic, and analgesic properties. In addition, by acting on AMPA (α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid) receptors, pregabalin indirectly reduces synaptic noradrenaline release (Fink 2002).

Notably, the mechanism of action of pregabalin does not appear to differ from that of gabapentin. The affinity of pregabalin for the α -2- δ modulatory site, however, is much greater than that of gabapentin. This explains why pregabalin is three- to six-fold more potent than gabapentin in animal models of seizures and epilepsy, and also in models of anxiety and neuropathic pain.

Why it is important to do this review

This review is an update of a previous Cochrane Review (Panebianco 2019), and aims to summarise existing data regarding the effects of add-on pregabalin for people with drug-resistant focal-onset seizures. Clinical trials published on the antiepileptic properties of pregabalin have so far focused on people with drug-resistant focal-onset epilepsy. In these randomised placebo-controlled trials, study participants are randomised to have either pregabalin or placebo added to their existing AED treatment. This is in keeping with international guidelines on the development of AEDs (ILAE Commission 1989). Once a drug has confirmed efficacy and safety as an add-on therapy, it can be tested as monotherapy. The use of pregabalin as monotherapy has been addressed in a separate Cochrane Review by Zhou 2012.

OBJECTIVES

To assess the efficacy and tolerability of pregabalin when used as an add-on treatment for drug-resistant focal epilepsy.

METHODS

Criteria for considering studies for this review

Types of studies

To be included in the review, studies had to meet the following criteria:

- 1. randomised controlled trials;
- 2. double-blind or single-blinded trials;
- 3. placebo controlled or active controlled;
- 4. parallel-group or cross-over studies.

Types of participants

We included people of any age with drug-resistant focal epilepsy (i.e. experiencing simple focal, complex focal, or secondary generalised tonic-clonic seizures).

Types of interventions

1. The active-treatment group received pregabalin in addition to an existing AED regimen taken at time of randomisation.

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2. The control group received a matched placebo or an active comparator control in addition to an existing AED regimen taken at time of randomisation.

Types of outcome measures

Primary outcomes

50% or greater reduction in seizure frequency

We chose the proportion of people with a 50% or greater reduction in seizure frequency in the treatment period compared to the pre-randomisation baseline period as the primary outcome. We chose this because it is a commonly reported outcome, and can be calculated for studies that do not report this outcome provided that baseline seizure data were recorded.

Secondary outcomes

Seizure freedom

We calculated this as the proportion of participants with a complete cessation of seizures during the treatment period.

Treatment withdrawal

We used the proportion of participants having treatment withdrawn for any reason during the course of the treatment period as a measure of global effectiveness. Treatment is likely to be withdrawn due to adverse effects, lack of efficacy, or a combination of both, and this is an outcome to which the individual makes a direct contribution. In trials of short duration, it is likely that adverse effects will be the most common reason for withdrawal. We also assessed the proportion of participants having treatment withdrawn for adverse effects.

Adverse effects

1. The proportion of participants experiencing the following five adverse effects (we considered these adverse effects to be common and important adverse effects of AEDs):

- a. ataxia (co-ordination problems);
- b. dizziness;
- c. fatigue;
- d. nausea;
- e. somnolence (unusual drowsiness).
- 2. The proportion of participants experiencing the five most common adverse effects mentioned in the included trials, if these differed from those listed as 1a to 1e. above.

Search methods for identification of studies

Electronic searches

For the latest update we searched the following databases on 16 November 2020:

- 1. Cochrane Register of Studies (CRS Web), using the search strategy outlined in Appendix 1. CRS Web includes randomised or quasi-randomised, controlled trials from PubMed, Embase, ClinicalTrials.gov, the World Health Organisation International Clinical Trials Registry Platform (ICTRP), the Cochrane Central Register of Controlled Trials (CENTRAL), and the Specialised Registers of Cochrane Review Groups, including Epilepsy.
- 2. MEDLINE (Ovid, 1946 to 16 November 2020) using the search strategy outlined in Appendix 2.

We did not impose any language restrictions.

Searches for this review were first run in 2007. The most recent electronic searches conducted in November 2020 identified a total of seven records that were potentially eligible for inclusion (see Figure 1). We automatically removed one record, because it was obviously irrelevant. We screened the titles and abstracts of the remaining six records, and excluded four records. We retrieved the full-text publications of the remaining two records, and we found them to be eligible for inclusion





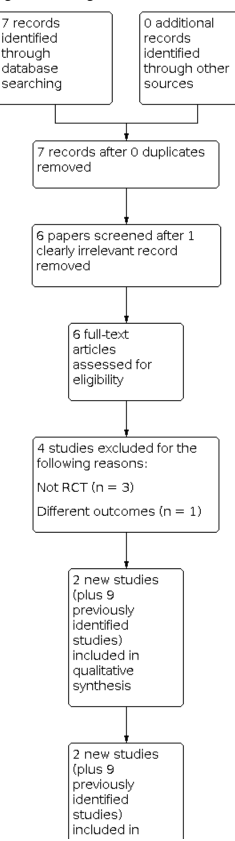




Figure 1. (Continued)

studies) included in quantitative synthesis (meta-analysis)

Searching other resources

We reviewed the reference lists of retrieved studies to check for additional reports of relevant studies. We also contacted Pfizer Ltd (manufacturer of pregabalin) and colleagues in the field.

Data collection and analysis

Selection of studies

For the update, two review authors (RB and MP) independently assessed trials for inclusion. Any disagreements were resolved by discussion with a third review author (AGM). Two review authors (RB and MP) extracted data and assessed risk of bias; again, disagreements were resolved by discussion.

Data extraction and management

The same two review authors (MP and RB) extracted the following information from the included trials, resolving any disagreements by mutual discussion.

Methodological/trial design

- 1. Method of randomisation and concealment.
- 2. Method of double-blinding.
- 3. Whether any participants had been excluded from the reported analyses.
- 4. Duration of baseline period.
- 5. Duration of treatment period.
- 6. Dose(s) of pregabalin tested.

Participant/demographic information

- 1. Total number of participants allocated to each treatment group.
- 2. Age/sex.
- 3. Number with focal/generalised epilepsy.
- 4. Seizure types.
- 5. Seizure frequency during the baseline period.
- 6. Number of background drugs.

For all trials sponsored by Pfizer Ltd, confirmation of the following information

- 1. Method of randomisation.
- 2. Total number randomised to each group.
- 3. Number of participants in each group achieving a 50% or greater reduction in seizure frequency.
- 4. Number of participants in each group having treatment withdrawn post randomisation.

- 5. For excluded participants:
 - a. the reason for exclusion;
 - b. whether any of those excluded completed the treatment phase;
 - c. whether any of those excluded had a 50% or greater reduction in seizure frequency during the treatment phase.

Outcomes

We recorded the number of participants experiencing each outcome per randomised group (see Types of outcome measures).

Assessment of risk of bias in included studies

For the update, two review authors (RB and MP) independently assessed risk of bias for each trial using the Cochrane risk of bias tool, as described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011), resolving any disagreements by discussion. We judged each included study to be at low, high, or unclear risk of bias for the six domains applicable to randomised controlled trials: randomisation method, allocation concealment, blinding methods, incomplete outcome data, selective outcome reporting, and other sources of bias.

Measures of treatment effect

We presented the primary outcome of seizure reduction as a risk ratio (RRs) and odds ratios (ORs) with 95% confidence intervals (CIs). We also presented the secondary outcomes, including seizure freedom, treatment withdrawal, and adverse effects, as risk ratios (RRs) and odds ratios (ORs) with 95% confidence intervals (CIs).

Unit of analysis issues

The inclusion of cross-over studies in meta-analyses introduces unit of analysis issues because each participant contributes data to both treatment groups. We had planned to extract data from the first treatment period of any eligible cross-over studies, had any been identified for inclusion. Essentially, we would have regarded the first treatment period as a parallel study, thus preventing data from the same participant being considered twice whilst simultaneously avoiding any issues of carry-over effect. We did not include any cross-over studies in this current review update, hence there were no unit of analysis issues to consider.

Dealing with missing data

We sought any missing data from the study authors. We carried out intention-to-treat (ITT), best-case, and worst-case analysis on the primary outcome to account for any missing data. All analyses are presented in the main report.

Assessment of heterogeneity

We assessed clinical heterogeneity by comparing the distribution of important participant factors amongst trials (e.g. age, seizure type, duration of epilepsy, number of antiepileptic drugs taken

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at the time of randomisation) and trial factors (e.g. allocation concealment, blinding, losses to follow-up). We examined statistical heterogeneity using a Chi² test and the I² statistic for heterogeneity and, providing no significant heterogeneity was present (P > 0.10), we employed a fixed-effect model. In the event that we found heterogeneity (P < 0.10), we used a random-effects model analysis employing the inverse-variance method.

Assessment of reporting biases

We requested protocols from study authors to enable a comparison of outcomes of interest. We investigated outcome reporting bias using the Outcome Reporting Bias in Trials (ORBIT I) matrix system for benefit outcomes (Kirkham 2010), and ORBIT II matrix system for harm outcomes (Saini 2014). We examined the asymmetry of funnel plots to detect any publication bias.

Data synthesis

We analysed the data using Review Manager (RevMan) version 5.4 (Review Manager 2020). Heterogeneity determined the choice of a fixed-effect or a random-effects model. If clinically appropriate, and if we found no evidence of substantial statistical heterogeneity using the I² statistic (I²< 50%), we analysed data in a meta-analysis using a fixed-effect model. If we found substantial heterogeneity (I² \geq 50%), we explored possible factors contributing to the heterogeneity and used a random-effects model to perform meta-analysis.

We carried out the following comparisons:

- 1. pregabalin intervention group versus placebo control group;
- 2. pregabalin intervention group versus active-comparator control group.

We planned to stratify each comparison by study characteristics, such as dose of pregabalin used, during subgroup analysis to ensure the appropriate combination of study data.

Our preferred estimator was the Mantel-Haenszel risk ratio (RR). For the outcomes 50% or greater reduction in seizure frequency and treatment withdrawal, we used 95% confidence intervals (CIs). For individual adverse effects, we used 99% CIs to make an allowance for multiple testing.

Our analyses included all participants in the treatment group to which they had been allocated. For the efficacy outcome (50% or greater reduction in seizure frequency), we undertook three analyses:

- 1. Primary (ITT) analysis: participants not completing follow-up or with inadequate seizure data were assumed to be non-responders. All of the included studies reported analysis by ITT.
- 2. Worst-case analysis: participants not completing follow-up or with inadequate seizure data were assumed to be nonresponders in the intervention group, and responders in the placebo group.
- 3. Best-case analysis: participants not completing follow-up or with inadequate seizure data were assumed to be responders in the intervention group, and non-responders in the placebo group.

The purpose of the best-case and worst-case analyses is to test whether the assumption made during ITT analysis (that all

participants not completing follow-up or with inadequate seizure data are non-responders) affects the estimated effect size.

Dose regression analysis

We undertook the dose-response analysis using a generalised linear mixed model (i.e. a model including both fixed and random effects) with the logit link function, as described in Turner 2000, and estimated using the command xtmelogit in STATA SE version 14 (Stata). We included the study and dose as fixed effects within the mixed model, whilst including treatment as a random effect within the mixed model (no random effect was included for the constant term of the mixed model). We standardised dose by its standard deviation (245 mg). This method estimated an odds ratio (OR) as opposed to a RR.

Subgroup analysis and investigation of heterogeneity

We undertook subgroup analysis for all the outcomes included in this review. For the comparison pregabalin versus placebo, subgroup analysis was stratified by dose of pregabalin. Dose of pregabalin was chosen because it was the most striking clinical difference identified between the included studies and was thus anticipated to be the cause of any observed heterogeneity. For the comparison pregabalin versus active comparator, subgroup analysis was stratified by active comparator in order to determine whether pregabalin might be advantageous or disadvantageous compared to a specific alternative AED. If deemed appropriate, we intended to investigate heterogeneity using sensitivity analysis.

Sensitivity analysis

We also intended to carry out sensitivity analysis if we found any peculiarities between studies' quality, characteristics of participants, interventions, and outcomes. We did not find any peculiarities between the studies, therefore we did not conduct any sensitivity analyses.

Summary of findings and assessment of the certainty of the evidence

We used the GRADE approach, as outlined in the GRADE Handbook (Schünemann 2013), to interpret findings, and GRADEpro GDT software (which imports data from Review Manager 5 software (GRADEpro GDT)), to create summary of findings tables for both comparisons: pregabalin versus placebo and pregabalin versus active comparator. We GRADE-assessed the following outcomes, deemed to be the most important: 50% or greater reduction in seizure frequency (intention-to-treat, best-case, and worst-case analysis), seizure freedom, treatment withdrawal for any reason, treatment withdrawal due to adverse effects and adverse effects (ataxia, dizziness, fatigue, headache, somnolence, weight gain). We assessed the evidence across eight criteria (risk of bias, inconsistency, indirectness, imprecision, publication bias, effect size, presence of plausible confounding factors, and dose-response gradient) to determine its certainty.

RESULTS

Description of studies

Results of the search

Searches for this review were first run in 2007. The most recent electronic searches conducted in November 2020 identified a total of seven records that were potentially eligible for inclusion (see Figure 1). We automatically removed one record, because it was obviously irrelevant. We screened the titles and abstracts of the remaining six records, and excluded four records. We retrieved the full-text publications of the remaining two records, and we found them to be eligible for inclusion.

Included studies

In this update we included two additional studies (Antinew 2019; Mann 2020). In the previous review, we included nine randomised controlled trials. The parallel-group studies in the review included people of any age (total 3949 participants). All 11 of the included studies recruited participants with drug-resistant focal-onset seizures. Participants were taking between one and four AEDs and had at least three, four, or six focal seizures per month in the pre-randomisation baseline period. All studies were sponsored by the drug manufacturer Pfizer Ltd.

We summarised the relevant information for each study in the Characteristics of included studies tables.

Antinew 2019 reported a multicentre trial (18 countries in Europe, Asia and the USA), comprising 295 participants. Randomised participants were between four and 16 years of age, receiving a stable regime of one to three AEDs. Treatment arms were: pregabalin 2.5 mg/kg/d (n = 104), pregabalin 10 mg/kg/d (n = 97), and placebo (n = 94). Maximum dose was 150 mg/d for the pregabalin 2.5 mg/kg/d (no dose escalation) arm, and 600 mg/d for the pregabalin 10 mg/kg/d arm (2.5 mg/kg/d for week one; 1.5 mg/kg/d for week two). This study had three phases: eight-week baseline, 12-week double-blind treatment (two-week dose escalation; 10-week fixed dose), and one-week taper. One participant randomised to the placebo group was excluded from the ITT analysis, because he or she was lost to follow-up.

Arroyo 2004 published a multicentre trial (45 sites in Europe, Australia, and Africa) that included 288 participants. Inclusion criteria were defined as people aged 18 years or older with focal-onset seizures. As an electroencephalogram (EEG) was not required to confirm the diagnosis, some of the 18 participants who were stated as having "generalised seizures", rather than secondary generalised, may have had primary generalised epilepsy. Participants were randomised to either 50 mg pregabalin three times daily (n = 99); 200 mg pregabalin three times daily (n = 92); or placebo three times daily (n = 97). After a baseline assessment of 8 weeks, the treatment period was conducted over 12 weeks with a 4 to 8 day titration period. During the treatment period, participants were assessed weekly for the first two weeks and fortnightly thereafter. Median follow-up was 12 weeks (range one day to 12 weeks). The study reported three time points, each at four-weekly intervals.

Baulac 2010 conducted a multicentre trial (97 sites in Europe, Canada, and Australia), comprising 434 participants. Randomised participants were between 16 and 82 years of age and had undergone an EEG within two years prior to randomisation. Treatment arms were 150 mg to 300 mg pregabalin twice daily (n = 152), 150 mg to 200 mg lamotrigine twice daily (n = 141), and placebo (n = 141). Following a six-week baseline period, there was a 17-week double-blind treatment period comprising two phases. The first phase (phase I) spanned 11 weeks and included an up-titration period (one week for pregabalin and five weeks for lamotrigine). During phase I, participants randomised to pregabalin and lamotrigine were both up-titrated to 300 mg/d of their respective treatment drugs and were then maintained on this dose. Participants who were seizure-free for the duration of phase I continued to be maintained on 300 mg/d active treatment for the duration of phase II (six weeks). Participants randomised to pregabalin who continued to have seizures were further up-titrated to 600 mg/d pregabalin during phase II, whereas participants randomised to lamotrigine who continued to have seizures were up-titrated to 400 mg/d lamotrigine for the remaining six weeks of the treatment period. The trial did not report participant review time points and follow-up.

Beydoun 2005 randomised 313 participants, aged 17 to 82 years, from 43 USA and Canadian centres in a randomised placebocontrolled trial. Treatment groups included 200 mg pregabalin three times daily (n = 111); 300 mg pregabalin twice daily (n = 104); and placebo (n = 98). After a baseline assessment of eight weeks, the treatment period was conducted over 12 weeks (including a one-week titration period). Follow-up occurred on weeks two, four, eight, and 12. Median follow-up was 12 weeks (range not reported). During the trial an interim analysis was carried out on 129 participants, which led to an alteration of the subsequent statistical analysis.

Elger 2005 reported a multicentre trial (53 sites in Canada and Europe) of 341 participants, aged 17 to 78 years. Treatment arms were 150 mg to 600 mg pregabalin (n = 131), titrated with respect to clinical response and adverse effects in 150 mg daily increments; fixed-dose pregabalin of 300 mg twice daily (n = 137); and placebo (n = 73). Participants were randomised to one of the three treatments using a 2:2:1 ratio, respectively. The treatment period ran over 12 weeks and followed a six-week baseline period. Participants were reviewed at two, four, eight, and 12 weeks into the study. Median follow-up was 12 weeks, and over 58% of participants completed the study in each arm. The trial did not report the range of follow-up.

French 2003 published a multicentre trial (76 sites in the USA and Canada) that included 455 participants. Randomised participants were between 12 and 70 years of age, but not all had EEG and imaging data. Participants were randomised into one of five treatment arms: 50 mg/day (n = 88), 150 mg/day (n = 88), 300 mg/day (n = 90), and 600 mg/day (n = 89) pregabalin in a twice-daily regimen, and placebo (n = 100). Baseline assessment occurred over eight weeks, and treatment duration was 12 weeks with no titration period. Follow-up occurred on weeks two, four, eight, and 12. Median follow-up was 12 weeks (range one day to 12 weeks). Around 83% of participants completed the study.

French 2014 published a multicentre trial (66 centres in the USA, Europe, and Asia) that included 325 participants, aged 18 to 75 years. Participants were randomised 1:1:1 to controlled-release pregabalin 165 mg (n = 101), 330 mg (n = 114), or placebo (n = 110). The trial ran over 23 weeks including an eight-week baseline phase, a two-week double-blind dose escalation, a 12-week double-blind maintenance phase, and a one-week taper. The mean overall compliance was 99.2% for all three treatment arms, as demonstrated by participant-completed diaries.

French 2016 conducted a multicentre trial (56 centres in Eastern and Western Europe, Asia, South and Central America) that included 484 participants between 18 and 80 years of age. Participants were randomised 1:1 to pregabalin 450 mg/d (n

Pregabalin add-on for drug-resistant focal epilepsy (Review) Copyright © 2022 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd. = 241) or gabapentin 1500 mg/d (n = 241). The trial included a six-week baseline phase (screening), a nine-week doubleblind dose escalation (titration) phase, and a 12-week doubleblind maintenance phase (21-week treatment phase overall). The primary endpoint was the percentage change in 28-day seizure rate from baseline to the treatment phase. Around 74% of participants completed the study.

Lee 2009 conducted a multicentre trial (nine sites in Korea) consisting of two treatment arms. A total of 178 participants, aged 18 years and above, were randomised to either 75 mg to 300 mg pregabalin twice daily (n = 119) or placebo (n = 59) using a 2:1 ratio. Following a six-week baseline period, treatment was conducted over 12 weeks with a one-week taper period at the end. Participants were assessed at weeks two, four, six, eight, and 12, with a follow-up visit at week 13. Eighty-eight per cent of randomised participants completed the study.

Mann 2020 reported a multicentre trial (22 countries in Europe, Asia, Russian Federation and the USA), comprising 175 participants. Inclusion criteria for this study included paediatric participants aged from one month up to four years old, with at least three seizures in the month prior to screening as observed by the parent/ caregivers and at least two seizures recorded during the 48- to 72hour baseline video-EEG monitoring. Participants were randomised to one of three treatment arms: pregabalin 7 mg/kg/d (n = 71), pregabalin 14 mg/kg/d (n = 34), and placebo (n = 70). This study had three phases: 48- to 72-hour baseline phase, 14-day doubleblind treatment phase (five-day dose escalation, 99-day fixed dose including video-EEG monitoring of 48 to 72 hours over final three days), and a seven-day double-blind taper phase.

Zaccara 2014 randomised 509 participants aged 18 years or older from 71 centres in Europe, the USA, and Asia to one of two

groups: pregabalin (n = 254; median dose 450 mg) or levetiracetam (n = 255; median dose 2000 mg). The trial included a six-week baseline phase, a four-week dose escalation phase, and a 12-week maintenance phase. During the trial an interim analysis was carried out after approximately 50% of participants had completed the maintenance phase. The trial continued as planned.

Excluded studies

We excluded four studies from this review update for the following reasons: three studies were not randomised controlled trials (Hu 2020; Morano 2019; Moseley 2019); one study described other types of outcomes measures (Aicua-Rapun 2020).

From the searches conducted since the previous review update, it was clear that the previously ongoing trial IRCT2012091210508N4 has since been published (Taghdiri 2015). However, we excluded this study due to ineligibility regarding the study population. Further details are provided in the Characteristics of excluded studies table.

Studies awaiting classification

We assessed two studies as awaiting classification (Russi 2006; Tata 2007), as we have obtained no additional information regarding either study since the publication of the previous review (see Characteristics of studies awaiting classification table).

Risk of bias in included studies

We assessed all 11 included studies for risk of bias based on the six domains of the Cochrane risk of bias tool. See Characteristics of included studies tables for each study for further details, Figure 2 for the risk of bias graph and Figure 3 for the risk of bias summary.

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.

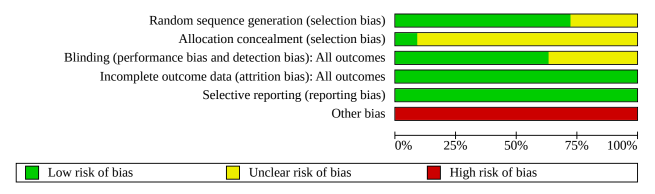
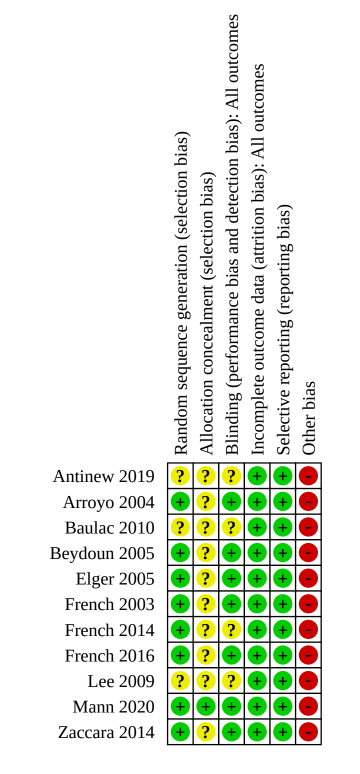




Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.





Allocation

For sequence generation, we rated eight studies at low risk of bias due to use of a computer-generated randomisation schedule or random permuted blocks (Arroyo 2004; Beydoun 2005; Elger 2005; French 2003; French 2014; French 2016; Mann 2020; Zaccara 2014). We rated three studies at unclear risk of bias due to lack of details of method use (Antinew 2019; Baulac 2010; Lee 2009).

In one trial, we rated the methods by which allocation was concealed at low risk of bias (Mann 2020). Ten trials did not provide information and were at unclear risk of bias for this domain (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; French 2014; French 2016; Lee 2009; Zaccara 2014).

Blinding

Seven studies were reported as double-blinded with the use of identical tablets with identical packaging for all treatment groups; we assessed these studies as at low risk of performance and detection bias (Arroyo 2004; Beydoun 2005; Elger 2005; French 2003; French 2016; Mann 2020; Zaccara 2014). No details were available for the other four studies, which we rated at unclear risk of performance and detection bias (Antinew 2019; Baulac 2010; French 2014; Lee 2009).

Incomplete outcome data

All studies reported study attrition rates, and all studies used an ITT analysis on randomised participants who took at least one dose of medication, using the 'last observation carried forward' approach, that is for participants failing to complete follow-up, seizure-frequency data were extrapolated from the last participant observation point for the whole treatment period, whilst for participants with no seizure data during the treatment period, baseline data were extrapolated.

Selective reporting

We requested the protocols for all included studies to compare a priori methods and outcomes against the published report, but these were unavailable. All studies reported the primary/secondary outcomes stated in the methods section in the results section of the articles. Notably, all expected outcomes with respect to this review were reported, therefore we had no suspicions or concerns about any purposefully withheld data. Further to this, we also completed an outcome matrix (see Figure 4) according to the ORBIT I and ORBIT II matrix system to investigate the potential for outcome reporting bias (Kirkham 2010; Saini 2014). Four included studies did not report seizure freedom (Arroyo 2004; French 2003; French 2014; French 2016), however we did not find this to be concerning. All of the participants in the included studies had drugresistant epilepsy, meaning that their epilepsy is refractory despite treatment with currently available antiepileptic medication. As a result, it is unlikely that many participants will achieve seizure freedom. Instead, 50% or greater reduction in seizure frequency is a more clinically relevant efficacy outcome for these participants. Consequently, it would be much more concerning and suspicious if any of the included studies failed to report the primary outcome, 50% or greater reduction in seizure frequency. In addition, we noted that several of the included studies did not report all of the adverse effects investigated as part of this review. However, because the majority of other harms were fully reported, we did not find this to be suspicious. The studies specified that they only reported the most common adverse events, for example only those reported by more than 5% of the study population, thus further justifying the absence of some data. We consequently rated all eleven included studies as at low risk of reporting bias (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; French 2014; French 2016; Lee 2009; Mann 2020; Zaccara 2014).



Figure 4. ORBIT Matrix

Study ID	Review primary outcome s	Review secondary outcomes					Revie	ew harm	outcomes		
(author, date of publicatio n)	50% or greater reductio n in seizure frequen cy	Seizure freedo m	Treatmen t withdraw al for any reason	Treatmen t withdraw al due to adverse events	Ataxi a	Dizzine ss	Fatigu e	Nause a	Somnolen ce	Headac he	Weig ht gain
Antinew 2019	1	×	4	4	×	×	~	×	1	1	~
Arroyo 2004	4	×	4	1	1	4	1	1	1	4	1
Baulac 2010	1	1	4	4	~	1	1	×	~	1	1
Beydoun 2005	4	-	4	1	1	1	1	1	4	×	1
Elger 2005	1	1	1	1	1	~	4	1	1	1	1
French 2003	1	×	1	1	1	1	1	×	1	4	1
French 2014	4	×	4	4	×	1	1	1	1	×	~
French 2016	4	×	-	4	×	1	×	×	4	1	1
Lee 2009	1	1	4	4	1	~	1	×	~	4	1
Mann 2020	~	×	4	~	×	×	×	×	~	×	×
Zaccara 2014	1	1	4	4	×	1	×	1	4	1	~

Other potential sources of bias

All studies were sponsored by Pfizer, the manufacturer of pregabalin; therefore, we rated all studies as having high risk of funding bias. In addition, in two included studies (Arroyo 2004; French 2003), individuals with primary generalised epilepsy may have been included in the trials, possibly leading to bias within the results.

Effects of interventions

See: Summary of findings 1 Pregabalin compared to placebo for drug-resistant focal epilepsy; Summary of findings 2 Pregabalin compared to active comparators for drug-resistant focal epilepsy

See Summary of findings 1 for the main comparison 'pregabalin versus placebo for drug-resistant epilepsy'.

Pregabalin versus placebo control

Eight included studies involving a total of 2338 randomised participants compared immediate-release pregabalin versus placebo (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; Lee 2009; Mann 2020). Another study (French 2014), including 325 participants, compared controlled-release pregabalin versus placebo. We included these nine trials in the analysis for the comparison 'pregabalin versus placebo'.

50% or greater reduction in seizure frequency

Nine included studies (2663 participants) reported this outcome (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; French 2014; Lee 2009; Mann 2020). An ITT analysis pooling all doses (50 mg to 600 mg/day immediate- and controlled-release pregabalin) showed evidence of heterogeneity ($I^2 = 77\%$), therefore we employed a random-effects model. Participants allocated to pregabalin were significantly more likely



to achieve a 50% or greater reduction in seizure frequency than those allocated to placebo (risk ratio (RR) 1.95, 95% confidence interval (CI) 1.40 to 2.72; Analysis 1.1). Subgroup analyses assessing the effect of individual doses showed no significant effect for 50 mg/d immediate-release pregabalin (RR 1.06, 95% CI 0.52 to 2.12; Analysis 2.1). Higher doses of immediate-release pregabalin were associated with a significantly higher proportion of participants achieving a 50% or greater reduction in seizure frequency compared to placebo (150 mg/d: RR 2.22, 95% CI 1.36 to 3.63; 300 mg/d: RR 2.86, 95% CI 1.65 to 4.94; 600 mg/d: RR 4.62, 95% CI 3.34 to 6.39; titrated 150 mg/d to 600 mg/d: RR 1.76, 95% CI 1.35 to 2.30; Analysis 2.1). Notably, during subgroup analysis, the effect size appeared to increase as the daily dose of pregabalin increased. Neither dose of controlled-release pregabalin, 165 mg/d or 330 mg/d, was associated with a significantly higher proportion of participants achieving a 50% or greater reduction in seizure frequency than placebo (165 mg/d: RR 1.03, 95% CI 0.72 to 1.48; 330 mg: RR 1.26, 95% CI 0.91 to 1.75; Analysis 2.1). Although the effect size did vary between subgroups, the direction of the effect (pregabalin being advantageous compared to placebo) was consistent amongst all subgroups, even those showing the smallest and statistically insignificant risk ratios.

Best-case and worst-case analyses

A best-case analysis (all treatment withdrawals in the treatment group assumed to be responders), pooling all doses (50 mg to 600 mg/day immediate- and controlled-release pregabalin), again showed that participants allocated to pregabalin were significantly more likely to achieve a 50% or greater reduction in seizure frequency compared to placebo 2.91 (RR 2.91, 95% CI 1.92 to 4.42; Analysis 1.2). Again, we detected significant heterogeneity within the data set ($I^2 = 87\%$) and so employed a random-effects model followed by a subgroup analysis according to dose to investigate dose as a potential source of heterogeneity (Analysis 2.2). Subgroup analyses assessing the effect of individual doses showed significant

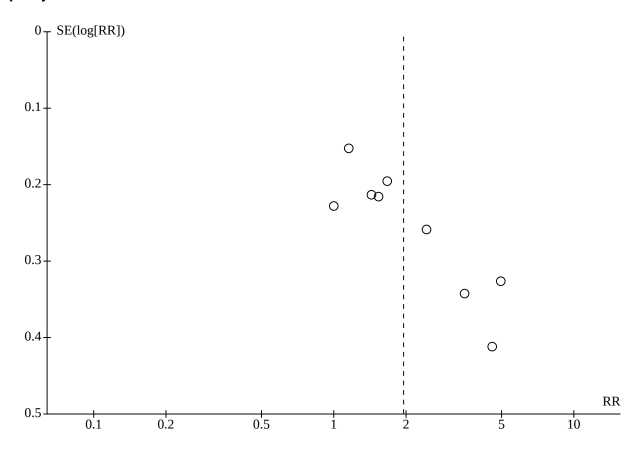
effects for all pregabalin doses, including 330 mg/d controlledrelease pregabalin (50 mg/d: RR 1.87, 95% CI 1.03 to 3.40; 150 mg/ d: RR 3.18, 95% CI 2.00 to 5.06; 300 mg/d: RR 4.37, 95% CI 2.61 to 7.29; 600 mg/d: RR 7.72, 95% CI 5.64 to 10.57; titrated 150 mg/d to 600 mg/d: RR 2.86, 95% CI 2.24 to 3.65; 330 mg/d controlledrelease: RR 1.63, 95% CI 1.21 to 2.20), with the exception of 165 mg/ d controlled-release pregabalin (RR 1.28, 95% CI 0.92 to 1.79), for which the effect size remained insignificant, even during best-case analysis (Analysis 2.2).

In contrast, a worst-case analysis (all dropouts from the control group assumed to be responders) pooling all doses of pregabalin (50 mg/d to 600 mg/d and including controlled-release pregabalin) showed no significant difference between pregabalin and placebo for the outcome 50% or greater seizure reduction (RR 1.10, 95%) CI 0.92 to 1.31; Analysis 1.3). The analysis did, however, continue to show significant heterogeneity within the data set ($I^2 = 56\%$). Subgroup analyses by dose indicated that one dose (50 mg/d) was associated with significantly fewer participants achieving a 50% or greater seizure reduction in the pregabalin group compared to the placebo group during worst-case analysis (RR 0.55, 95% CI 0.30 to 0.99). In contrast, 600 mg/d pregabalin continued to demonstrate a significant advantage over placebo with respect to the number of participants who achieved a 50% or greater reduction in seizure frequency (RR 1.72, 95% CI 1.42 to 2.09). There were no significant differences between the other pregabalin dose groups and placebo for the outcome during worst-case analysis (150 mg/d: RR 0.96, 95% CI 0.66 to 1.38; 300 mg/d: RR 1.48, 95% CI 0.98 to 2.23; titrated 150 mg/d to 600 mg/d: RR 0.92, 95% CI 0.76 to 1.12; 165 mg/d controlled-release: RR 0.79, 95% CI 0.57 to 1.09; 330 mg/d controlled-release: RR 0.96, 95% CI 0.72 to 1.28; Analysis 2.3).

Furthermore, the ITT and best-case analyses had to been downgraded once more because we strongly suspected publication bias, as demonstrated by the funnel plot generated (Figure 5).



Figure 5. Funnel plot of comparison: 1 Pregabalin versus placebo, outcome: 1.1 50% or greater reduction in seizure frequency - ITT.



Dose regression analysis for 50% response

We fitted a generalised linear mixed model to the data from Analysis 1.1 (Figure 5) to estimate the effect of dose on the primary outcome, 50% or greater reduction in seizure frequency (details in Data synthesis). This method estimates an odds ratio (OR) as opposed to an RR. Dose was standardised by its standard deviation (245 mg). The odds of response (50% reduction in seizure frequency) approximately doubled (OR 1.99, 95% CI 1.74 to 2.28) with estimated between-study standard deviation of 0.17 (standard error 0.13) for each 245 mg increase in dose of pregabalin. This translates into an estimated doubling of odds of response with an increase in dose of 245 mg (e.g. a doubling of odds from approximately 300 mg to 600 mg).

Seizure freedom

Four included studies involving a total of 1125 participants reported seizure freedom (Baulac 2010; Beydoun 2005; Elger 2005; Lee 2009). Specifically, these studies reported the number of participants who had complete cessation of their seizures over the entire treatment period. In contrast, the studies by Antinew 2019 and Arroyo 2004 recorded the number of participants who were seizure-free during the last 28 days of their treatment. This definition of seizure freedom was not consistent with the definition used by the other studies, therefore we excluded the data extracted from Antinew 2019 and Arroyo 2004 from the analysis. The pooled analysis, consisting of all doses, showed evidence of no heterogeneity ($I^2 = 11\%$), therefore we continued to employ a fixed-effect model. The

analysis demonstrated that participants allocated to pregabalin were significantly more likely to attain seizure freedom than those allocated to placebo (RR 3.94, 95% CI 1.50 to 10.37; Analysis 1.4).

Although we detected no important heterogeneity, we continued to use subgroup analysis to investigate whether there was any potentially undetected heterogeneity due to experimental dose of pregabalin. Two subgroups were included in the analysis, 600 mg/ d pregabalin and 150 mg/d to 600 mg/d titrated dose of pregabalin. Subgroup analysis highlighted that a significantly greater number of participants randomised to 600 mg/d pregabalin attained seizure freedom compared to those randomised to placebo (RR 6.92, 95% CI 1.31 to 36.70; Analysis 2.4). There was no significant difference in the proportion of participants who achieved seizure freedom between participants allocated to 150 mg/d to 600 mg/d titrated dose of pregabalin compared to placebo (RR 2.39, 95% CI 0.83 to 6.89; Analysis 2.4).

Treatment withdrawal for any reason

Nine included studies (2663 participants) reported this outcome (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; French 2014; Lee 2009; Mann 2020). An analysis pooling all doses (50 mg/d to 600 mg/d immediate- and controlled-release pregabalin) showed no evidence of heterogeneity ($I^2 = 0\%$), therefore we used a fixed-effect model. Participants allocated to pregabalin were significantly more likely to have withdrawn from treatment compared to those allocated to placebo (RR 1.33, 95% CI 1.10 to 1.60; Analysis 1.5). Subgroup analysis assessing the

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individual doses showed no significant effect for 50 mg/d (RR 0.87, 95% CI 0.40 to 1.89); 150 mg/d (RR 0.72, 95% CI 0.41 to 1.28); 300 mg/ d (RR 1.62, 95% CI 0.85 to 3.10); or 150 mg/d to 600 mg/d titrated dose of immediate-release pregabalin (RR 1.20, 95% CI 0.89 to 1.62) compared to placebo. Similarly, neither dose of controlled-release pregabalin displayed a significant effect on the rate of treatment withdrawal (165 mg/d controlled-release: RR 0.82, 95% CI 0.36 to 1.86; 330 mg/d controlled-release: RR 1.21, 95% CI 0.59 to 2.46). The only dose of pregabalin associated with a significantly higher treatment withdrawal rate for any reason compared to placebo was 600 mg/d (RR 1.84, 95% CI 1.42 to 2.40; Analysis 2.5). In fact, the two lowest doses of immediate-release pregabalin (50 mg/d and 150 mg/d) and lowest dose of controlled-release pregabalin (165 mg/ d) both actually estimated risk ratios of less than one (50 mg/d: RR 0.87, 95% CI 0.40 to 1.89; 150 mg/d: RR 0.72, 95% CI 0.41 to 1.28; 165 mg/d controlled-release: RR 0.82, 95% CI 0.36 to 1.86), meaning that fewer people randomised to low doses of pregabalin withdrew from treatment compared to those randomised to placebo.

Treatment withdrawal due to adverse effects

Nine included studies (2663 participants) reported this outcome (Antinew 2019; Arroyo 2004; Baulac 2010; Beydoun 2005; Elger 2005; French 2003; French 2014; Lee 2009; Mann 2020). An analysis pooling all doses (50 mg/d to 600 mg/d immediate- and controlledrelease pregabalin) showed no evidence of heterogeneity ($I^2 = 0\%$), thus we used a fixed-effect model for the analysis. Participants allocated to pregabalin were significantly more likely to withdraw from treatment due to adverse effects (RR 2.60, 95% CI 1.86 to 3.64; Analysis 1.6). Subgroup analyses assessing treatment withdrawal with differing doses suggested that a higher withdrawal rate was associated with higher doses of pregabalin (50 mg/d: RR 1.36, 95% CI 0.43 to 4.31; 150 mg/d: RR 1.02, 95% CI 0.45 to 2.32; 300 mg/ d: RR 2.89, 95% CI 1.07 to 7.78; 600 mg/d: RR 3.78, 95% CI 2.47 to 5.81; Analysis 2.6). Specifically, participants randomised to 300 mg/d, 600 mg/d, or a titrated dose of pregabalin of 150 mg/d to 600 mg/d (RR 2.26, 95% CI 1.30 to 3.95) were all significantly more likely to withdraw from treatment due to adverse effects than were participants randomised to placebo. Neither dose of controlledrelease pregabalin was associated with a significantly different treatment withdrawal rate due to adverse effects compared to placebo (165 mg/d controlled-release: RR 1.09, 95% CI 0.22 to 5.27; 330 mg/d controlled-release: RR 2.57, 95% CI 0.70 to 9.45).

Adverse effects

In addition to the five prespecified adverse effects, weight gain and headache were among the most common adverse effects reported. Analyses pooling across doses (50 mg/d to 600 mg/d immediateand controlled-release pregabalin) indicated that ataxia (RR 3.90, 99% CI 2.05 to 7.42; Analysis 1.7); dizziness (RR 3.15, 99% CI 2.23 to 4.44; Analysis 1.8); fatigue (RR 1.35, 99% CI 0.94 to 1.93; Analysis 1.9); somnolence (RR 2.05, 99% CI 1.49 to 2.81; Analysis 1.12); and weight gain (RR 4.35, 99% CI 2.34 to 8.11; Analysis 1.13) were all significantly more prevalent in participants randomised to pregabalin compared to placebo. Nausea incidence did not differ significantly between pregabalin and placebo groups (RR 1.20, 99% CI 0.56 to 2.58; Analysis 1.11). In contrast, participants randomised to pregabalin were significantly less likely to experience headache compared to those randomised to placebo (RR 0.65, 99% CI 0.45 to 0.94; Analysis 1.10). We detected no significant heterogeneity for any of the adverse effects analysed ($I^2 = 0\%$).

Subgroup analysis according to dose of pregabalin revealed that the highest dose, 600 mg/d pregabalin, was consistently associated with a significantly greater likelihood of participants experiencing adverse effects compared to placebo. Specifically, participants receiving 600 mg/d were more likely to experience the following adverse effects than participants receiving placebo: ataxia (RR 4.49, 99% CI 2.25 to 8.95; Analysis 2.7); dizziness (RR 3.72, 99% CI 2.42 to 5.69; Analysis 2.8); somnolence (RR 2.57, 99% CI 1.64 to 4.03; Analysis 2.12); and weight gain (RR 5.88, 99% CI 2.52 to 13.73; Analysis 2.13). Similarly, a titrated dose of 150 mg/d to 600 mg/d pregabalin was associated with a significantly increased incidence rate of ataxia (RR 4.46, 99% CI 1.28 to 15.48; Analysis 2.7); dizziness (RR 3.08, 99% CI 1.80 to 5.28; Analysis 2.8); somnolence (RR 2.35, 99% CI 1.31 to 4.19; Analysis 2.12); and weight gain (RR 3.64, 99% CI 1.49 to 8.87; Analysis 2.13). In contrast, none of the other dose subgroups were consistently associated with an increased likelihood of the individual adverse effects. Interestingly, all the dose groups, except for 50 mg/d pregabalin (RR 1.01, 99% CI 0.31 to 3.33), were, however, associated with an increased incidence of dizziness compared to placebo (Analysis 2.8). Furthermore, all the dose subgroups had a risk ratio suggesting that there was a decreased likelihood of participants experiencing headache when receiving pregabalin compared to those receiving placebo, however the difference was only significant for one subgroup, 150 mg/d pregabalin (RR 0.53, 99% CI 0.24 to 1.17; Analysis 2.10).

Pregabalin versus active comparator

Three included studies involving a total of 1286 participants compared pregabalin to other existing AEDs as active comparators. One study included 293 randomised participants and compared pregabalin with lamotrigine as the active control drug (Baulac 2010). Another trial included 509 randomised participants and compared pregabalin with levetiracetam as the active control drug (Zaccara 2014). The remaining study, French 2016, involved 484 participants and compared pregabalin with gabapentin as the active control drug. Summary of findings 2 presents the data for pregabalin versus active comparators.

50% or greater reduction in seizure frequency

All three included studies (1286 participants) reported this outcome. We detected significant heterogeneity within the data set $(I^2 = 61\%)$, therefore we used a random-effects model. The likelihood of participants achieving a 50% or greater reduction in seizure frequency was not significantly different based on whether participants were randomised to pregabalin or an alternative active-comparator AED (RR 1.03, 95% CI 0.85 to 1.25; Analysis 3.1). Interestingly, subgroup analysis according to active-comparator control group revealed that participants receiving pregabalin were significantly more likely to achieve a 50% or greater reduction in seizure frequency compared to those receiving the active comparator lamotrigine (RR 1.47, 95% CI 1.03 to 2.12; Analysis 4.1), however there was no significant difference between pregabalin and either levetiracetam (RR 0.94, 95% CI 0.80 to 1.11) or gabapentin (RR 0.96, 95% CI 0.82 to 1.12). The test for subgroup differences did not highlight a significant subgroup effect (P = 0.07; Analysis 4.1).

Best-case and worst-case analyses

We subsequently conducted a best-case analysis (all dropouts assumed to be responders to treatment). This revealed a significant increase in the proportion of participants who achieved a 50%

or greater seizure reduction in favour of the pregabalin group compared to active comparators (RR 1.60, 95% CI 1.17 to 2.19; Analysis 3.2). However, a worst-case analysis (all dropouts assumed to be responders to control) revealed a significant increase in the proportion of participants who achieved a 50% or greater seizure reduction in favour of the control group (RR 0.67, 95% CI 0.62 to 0.74; Analysis 3.3).

This was similarly the case when best-case and worst-case analysis was performed for each of the individual active comparators during subgroup analysis. Pregabalin appeared to be more efficacious than lamotrigine (RR 2.73, 95% CI 1.99 to 3.74); levetiracetam (RR 1.27, 95% CI 1.11 to 1.46); and gabapentin (RR 1.34, 95% CI 1.18 to 1.52) during best-case analysis (Analysis 4.2), but was shown to perform significantly worse than the active comparators during worst-case analysis (pregabalin versus lamotrigine: RR 0.68, 95% CI 0.52 to 0.88; pregabalin versus levetiracetam: RR 0.71, 95% CI 0.62 to 0.82; pregabalin versus gabapentin: RR 0.64, 95% CI 0.57 to 0.73; Analysis 4.3). The test for subgroup differences detected significantly different effect sizes between the subgroups during best-case scenario analysis (P < 0.001; Analysis 4.2), but not during worst-case analysis (P = 0.57; Analysis 4.3).

Seizure freedom

Only two included studies involving a total of 802 participants reported this outcome (Baulac 2010; Zaccara 2014). French 2016 reported no data regarding seizure freedom for this comparison. Participants randomised to pregabalin were significantly less likely to attain seizure freedom than participants randomised to an active comparator (RR 0.59, 95% CI 0.37 to 0.95; Analysis 3.4). When analysed separately during subgroup analysis according to activecomparator control group, the proportion of participants attaining seizure freedom was not significantly different for those receiving pregabalin compared to those receiving the active comparator lamotrigine (RR 1.39, 95% CI 0.40 to 4.83; Analysis 4.4). However, the seizure freedom rate was significantly lower in participants receiving pregabalin than in those receiving levetiracetam (RR 0.50, 95% CI 0.30 to 0.85; Analysis 4.4). Despite this, the test for subgroup differences did not reveal any significant differences between the effect size estimated by the two active-comparator control groups (P = 0.14; Analysis 4.4).

Treatment withdrawal for any reason

Three included studies involving a total of 1286 randomised participants reported this outcome (Baulac 2010; French 2016; Zaccara 2014). We found no significant difference in the rate of treatment withdrawal for any reason between pregabalin and active comparators (RR 0.93, 95% CI 0.76 to 1.13; Analysis 3.5). Similarly, during subgroup analysis according to active comparator, pregabalin was not shown to have a significantly different rate of treatment withdrawal for any reason compared to any of the individual active comparators: lamotrigine (RR 1.07, 95% CI 0.75 to 1.52); levetiracetam (RR 1.03, 95% CI 0.71 to 1.49); and gabapentin (RR 0.78, 95% CI 0.57 to 1.07) (Analysis 4.5). Furthermore, the test for subgroup differences was not statistically significant (P = 0.36).

Treatment withdrawal due to adverse effects

Three included studies involving a total of 1286 randomised participants also reported treatment withdrawal due specifically to adverse effects experienced (Baulac 2010; French 2016; Zaccara 2014). Again, there was no significant difference in the proportion of

participants who withdrew from treatment due to adverse effects between those randomised to pregabalin compared to an active comparator (RR 1.04, 95% CI 0.73 to 1.48; Analysis 3.6). According to the subgroup analysis, pregabalin did not demonstrate a significantly different treatment withdrawal rate due to adverse effects compared to any of the individual active comparators: lamotrigine (RR 0.89, 95% CI 0.53 to 1.48); levetiracetam (RR 1.29, 95% CI 0.66 to 2.54); or gabapentin (RR 1.07, 95% CI 0.54 to 2.11) (Analysis 4.6). Again, the test for subgroup difference was statistically insignificant (P = 0.69).

Adverse effects

Although the three included studies involving a total of 1286 participants documented and reported adverse effects, they did not report all the adverse effects we were investigating in this review (Baulac 2010; French 2016; Zaccara 2014). Notably, for some adverse effects, data were only supplied by one study, namely for ataxia, fatigue, and nausea. Furthermore, we detected significant heterogeneity within the data sets for the following adverse effects: dizziness ($I^2 = 65\%$), headache ($I^2 = 64\%$), and weight gain ($I^2 = 60\%$). We therefore used a random-effects model for the analysis of these adverse effects.

The occurrence of ataxia (RR 1.72, 99% CI 0.54 to 5.55; Analysis 3.7); fatigue (RR 1.72, 99% CI 0.77 to 3.83; Analysis 3.9); headache (RR 0.83, 99% CI 0.41 to 1.65; Analysis 3.10); and somnolence (RR 1.16, 99% CI 0.88 to 1.53; Analysis 3.12) did not differ significantly between pregabalin and active-comparator treatment groups. More participants randomised to pregabalin compared to those randomised to active comparators experienced dizziness (RR 1.64, 99% CI 0.85 to 3.16; Analysis 3.8) and weight gain (RR 2.87, 99% CI 0.94 to 8.75; Analysis 3.13). In contrast, significantly fewer participants randomised to pregabalin compared to those randomised to active comparators experienced nausea (RR 0.20, 99% CI 0.04 to 1.01; Analysis 3.11). Importantly, however, only one study, Zaccara 2014, provided data for this outcome, so it is difficult to draw any conclusions from this result.

Subgroup analysis according to active-comparator control group revealed that pregabalin was associated with a higher incidence rate for some of the adverse effects investigated compared to two of the active-comparator controls, lamotrigine and levetiracetam. Specifically, participants receiving pregabalin were more likely than those receiving the active comparator lamotrigine to experience the following adverse effects: dizziness (RR 2.94, 99% CI 1.32 to 6.52; Analysis 4.7); somnolence (RR 1.99, 99% CI 0.91 to 4.33; Analysis 4.10); and weight gain (RR 4.33, 99% CI 0.86 to 21.68; Analysis 4.11). Similarly, participants receiving pregabalin were more likely than those receiving the active comparator levetiracetam to experience dizziness (RR 1.44, 99% CI 0.89 to 2.34; Analysis 4.7) and weight gain (RR 4.82, 99% CI 1.39 to 16.74; Analysis 4.11). Interestingly, the incidence rate of adverse effects for pregabalin and gabapentin was not significantly different. Notably, however, pregabalin was associated with a significantly lower rate of headache compared to lamotrigine (RR 0.52, 99% CI 0.26 to 1.05; Analysis 4.8), as well as a significantly lower rate of nausea compared to levetiracetam (RR 0.20, 99% CI 0.04 to 1.01; Analysis 4.9), as alluded to earlier. Importantly, only one study provided data to each of the subgroups included in the subgroup analysis. Interestingly, the test for subgroup differences for each of the individual adverse effects indicated that there was not a significant subgroup effect

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dependent on active-comparator control (dizziness: P = 0.06; headache: P = 0.06; somnolence: P = 0.15; weight gain: P = 0.08).

DISCUSSION

Summary of main results

We identified eight randomised placebo-controlled parallel trials (Antinew 2019; Arroyo 2004; Beydoun 2005; Elger 2005; French 2003; French 2014; Lee 2009; Mann 2020), two active-comparatorcontrolled parallel trials (French 2016; Zaccara 2014), and one trial that included both an active-drug group and a placebo control group (Baulac 2010). All eleven studies were industry-sponsored (Pfizer Ltd). We took summary trial data from the relevant publications, and did not obtain any individual participant data. We attempted to retrieve the respective trial protocols but were unsuccessful. All studies appeared to be of good methodological quality overall. Most were randomised using suitable sequence generation methods, however only two studies reported their methods for concealing allocation. All included studies were reported to be double-blind, but only six studies provided adequate details of how blinding was achieved and maintained. Few participants were lost to follow-up. However, overall attrition rates were high for certain studies, especially those that included higher doses of pregabalin.

The included studies tested doses of pregabalin ranging from 50 mg/d to 600 mg/d, including both immediate- and controlledrelease pregabalin. The results showed that pregabalin, when used as an add-on treatment, can reduce seizure frequency in individuals with drug-resistant focal epilepsy. In the main analysis, when all doses of pregabalin were pooled, the RR for a 50% or greater reduction in seizure frequency was 1.95 (95% CI 1.40 to 2.72). A summary of the main findings for the pooled analysis of all doses of pregabalin versus placebo can be found in the Summary of findings 1.

We detected significant heterogeneity within the data set for the outcome 50% or greater reduction in seizure frequency. Whilst all doses of immediate-release pregabalin above 50 mg/d were observed to significantly increase 50% responder rate (the proportion of participants achieving 50% or greater seizure reduction) during ITT analysis, only 600 mg/d pregabalin consistently showed a significant therapeutic effect compared to placebo during ITT, best-case, and worst-case analysis. This suggests that the effect noted at 600 mg/d is a true effect, whereas it is possible that the therapeutic effect reported at the lower doses of pregabalin could be as a result of treatment withdrawals and may, therefore, not be accurate of the true efficacy of pregabalin. Furthermore, the significant therapeutic effect (an increased proportion of participants achieving 50% or greater seizure reduction) observed during pooled analysis was not consistently detected. Specifically, it was not reported during worst-case analysis. This raises doubts about the validity of the pooled effect described, and suggests that the therapeutic effect may be dose-dependent and might only be observed at the higher doses of pregabalin, rather than being a feature of all doses of pregabalin, generally.

The alternative efficacy outcome, seizure freedom, again emphasised the therapeutic potential of pregabalin. Participants allocated to pregabalin were significantly more likely to attain seizure freedom. However, data for this outcome were only provided by four studies, which accounted for two subgroups, 600 mg/d pregabalin and titrated dose of 150 mg/d to 600 mg/d. Consequently, we have no data specifically for the lower doses of pregabalin that were included in the previous efficacy analyses for 50% or greater seizure reduction. This potentially explains the lack of heterogeneity present in the seizure freedom data set compared to that revealed for the primary efficacy outcome. Again, the subgroup analysis indicated a high risk ratio for 600 mg/d (RR 6.92, 95% CI 1.31 to 36.70), emphasising a large treatment effect at this dose. In this scenario, it is estimated that if 1000 people were to receive pregabalin, 41 would likely achieve complete cessation of seizures, compared to 6 people if 1000 people were to receive placebo. Notably, other than 50% or greater reduction in seizure frequency, none of the other outcomes for the comparison pregabalin versus placebo displayed significant heterogeneity.

Although both efficacy outcomes recognised 600 mg/d pregabalin as a highly efficacious dose, it is important to acknowledge that 600 mg/d pregabalin was also associated with tolerability issues. This was the only dose to display a significantly higher withdrawal rate for any reason compared to placebo during subgroup analysis, and was one of only three doses to display a significantly higher treatment withdrawal rate due specifically to adverse effects. Notably, 600 mg/d pregabalin demonstrated the greatest risk ratio (RR 3.78, 95% CI 2.47 to 5.81) for treatment withdrawal due to adverse effects, indicating a very large effect size. Accordingly, 600 mg/d pregabalin was repeatedly associated with an increased incidence rate for the majority of the adverse effects investigated, namely ataxia, dizziness, fatigue, somnolence, headache, nausea and weight gain.

When pregabalin was compared to other AEDs, rather than placebo, it did not show a significant therapeutic advantage with regard to 50% responder rate during pooled analysis. During subgroup analysis, however, pregabalin was associated with a significantly higher responder rate compared to lamotrigine, but not when compared to levetiracetam or gabapentin. Surprisingly, the pooled analysis for the alternative efficacy outcome showed that participants randomised to pregabalin were significantly less likely to attain seizure freedom. Although we did not detect statistical heterogeneity in the data set, the two studies (each contributing to one subgroup) indicated opposing effects. The study comparing pregabalin to lamotrigine estimated an insignificant therapeutic effect (RR 1.39, 95% CI 0.40 to 4.83), whereas the study comparing pregabalin to levetiracetam indicated a diminished seizure freedom rate for participants randomised to pregabalin (RR 0.50, 95% CI 0.30 to 0.85). We detected no significant differences in treatment withdrawal rate for any reason or due to adverse effects, specifically, during either pooled analysis or subgroup analysis.

The majority of the adverse effects investigated, namely ataxia, fatigue, dizziness, headache, somnolence, nausea and weight gain were no more prevalent in participants randomised to pregabalin than in those randomised to active control. Participants receiving pregabalin did appear to be much more likely to experience weight gain compared to participants receiving active control. Specifically, during subgroup analysis large effect sizes were recognised for pregabalin versus both lamotrigine and levetiracetam. Notably, however, subgroup analysis did not reveal any differences in incidence rates of individual adverse effects between pregabalin and gabapentin. This is not surprising given that pregabalin and gabapentin are structurally related. Both drugs are structural analogues of γ -aminobutyric acid (GABA) and both bind with high affinity to the α -2- δ subunit of voltage-gated calcium channels (Bockbrader 2010).

Overall completeness and applicability of evidence

Heterogeneity was a serious issue for the outcome 50% or greater reduction in seizure frequency for both comparisons, pregabalin versus placebo and pregabalin versus active comparator. As a result, the treatment effect described and estimated for each pooled analysis may only be minimally informative. For the comparison pregabalin versus placebo, we combined a large range of doses, including a titrated dose regimen, into a single meta-analysis. It is less clear what dose individual participants were actually receiving during a titrated dose regimen. Data for many of these participants could have been entered into specific dose subgroups (i.e. 150 mg/d, 300 mg/d, or 600 mg/d) if the stratified data to enable this had been available in the relevant trial publications. This further complicates the meta-analysis.

Additionally, the pooled analysis for the comparison pregabalin versus placebo included both immediate-release and controlledrelease pregabalin, which have very different pharmacokinetics. This explains the difference in effect size calculated and the apparent heterogeneity between subgroups. For the other comparison, pregabalin versus active comparator, it is possible that the other AEDs equally have very different mechanisms of actions and potencies, therefore, it is difficult to combine them into a meta-analysis. As a result, for both comparisons, the pooled effect is unlikely to be representative of what will occur at every dose of pregabalin, or to reflect what the anticipated effect size of pregabalin is compared to another AED.

For this reason, the effect sizes reported from the subgroup analyses should be considered more informative than the effect size reported from the pooled analysis for the outcome 50% or greater reduction in seizure frequency. The variation in the effect reported thus limits our ability to sufficiently answer the question of whether pregabalin is more efficacious than placebo when used as an add-on treatment for drug-resistant focal epilepsy. The overall consensus appears to be that higher doses of immediate-release pregabalin, specifically 150 mg/d and greater, are more efficacious than placebo with regard to the 50% responder rate. However, pregabalin does not appear to offer a competitive advantage over other AEDs.

With further regard to subgroup analysis, it is important to recognise that for each active-comparator subgroup (i.e. pregabalin versus lamotrigine, pregabalin versus levetiracetam, and pregabalin versus gabapentin) for the comparison pregabalin versus active comparator, data were only supplied by one study, therefore were very limited. Similarly, for the comparison pregabalin versus placebo, data were only provided by one study for four of the subgroups: 50 mg/d, 300 mg/d, 165 mg/d controlledrelease, and 330 mg/d controlled-release pregabalin. As a result, multiple subgroups may have been underpowered and therefore any conclusions reached must be interpreted cautiously.

An additional issue is that the data included in this review were mainly derived from adult study populations (participants aged 16 years and above). Only three studies included a subset of younger participants (Antinew 2019; French 2003; Mann 2020).

Consequently, the findings reported in this review are only applicable to adults and are not informative of the effects of pregabalin in children.

Quality of the evidence

We assessed the certainty of the evidence for the primary and secondary outcomes for the comparison pregabalin versus placebo and pregabalin versus active comparator using the GRADE approach. The GRADE assessment is presented and summarised in Summary of findings 1. Overall, we rated the evidence as very low to moderate in certainty. We downgraded all outcomes once due to the unclear risk of bias across studies, mainly because all of the included studies failed to describe how allocation was concealed. Three of these studies also did not provide details about either the generation of the randomisation sequence or how blinding was effectively achieved.

We further downgraded the certainty of evidence for all three analyses, ITT, best-case, and worst-case analysis, for the outcome 50% or greater reduction in seizure frequency due to the significant statistical heterogeneity detected. As explained above, statistical heterogeneity greatly affected our ability to answer our hypothesis and impacted the validity of our conclusion. Furthermore, the ITT and best-case analyses had to been downgraded once more because we strongly suspected publication bias, as demonstrated by the funnel plot generated (Figure 6). It is clear from the funnel plot that the larger studies are predicting a much smaller effect size than the smaller studies. Ideally, we would expect the data points to produce a funnel shape, with the risk ratios estimated by the individual studies evenly distributed either side of the estimated pooled effect. Instead, the data points plotted from the individual study effects look more similar to a linear regression, suggesting publication bias.

Neither of the outcomes concerning treatment withdrawal or the other efficacy outcome, seizure freedom, were affected by either heterogeneity or suspected publication bias. Both seizure freedom and treatment withdrawal due to adverse effects were, however, rarer events compared to the outcome 50% or greater reduction in seizure frequency, and therefore, the number of events reported did not satisfy the optimal information size necessary for a robust meta-analysis. As a consequence, we downgraded the certainty of the evidence to low. Nevertheless, we were able to then upgrade the certainty of the evidence back to moderate for the outcomes seizure freedom and treatment withdrawal due to adverse effects because of the large effect size noted for each (RR > 2.00). We also upgraded the certainty of the evidence for the outcome 50% or greater reduction in seizure frequency for both the ITT and bestcase analysis, under the same principle. We further upgraded the certainty of the evidence for the ITT analysis as a result of the dose-response relationship detected by the regression analysis. This produced an overall judgement of low certainty of evidence for the primary efficacy outcome.

The low certainty of evidence for the outcome 50% or greater reduction in seizure frequency means that we are uncertain whether the effect size estimated is accurate of the true efficacy of pregabalin. In contrast, the rating of moderate certainty of evidence for the other three outcomes, seizure freedom and treatment withdrawal for any reason and due to adverse effects, means that we are fairly certain that the effect size reported is an accurate estimate of the true effect size.



Potential biases in the review process

The approach to analysis for all of the included trials used the 'last observation carried forward' method. For participants failing to complete follow-up, seizure frequency data were extrapolated to the whole treatment period, whereas for participants with no seizure data during the treatment period, baseline data were extrapolated. Whilst this approach may help minimise bias due to losses to follow-up (and is preferred by drug regulatory authorities), its use must be taken into consideration when interpreting the results of this systematic review, especially due to the high attrition rate noted in certain studies.

Importantly, the 'last observation carried forward' method assumes that a participant's response does not alter after treatment withdrawal. Specifically, the method does not consider any fluctuations in a participant's response or incorporate any imputation uncertainty. Consequently, the method likely predicts narrower confidence intervals than would normally be observed and, as a result, the effect size estimate is more likely to be statistically significant. Care must therefore be taken when considering the significance of the results presented.

Agreements and disagreements with other studies or reviews

We were able to identify two other systematic reviews that investigated pregabalin as an add-on therapy for drug-resistant focal epilepsy. Neither review included a meta-analysis, but both emphasised the therapeutic potential of pregabalin compared to placebo. Specifically, both reviews reported the outcome 50% or greater reduction in seizure frequency as evidence of pregabalin's antiepileptic effect. One review reported a responder rate (the proportion of participants achieving a 50% or greater reduction in seizure frequency) of 31% to 51% (Hamandi 2006), whilst the other reported a responder rate of 14% to 51% (Ryvlin 2008). Notably, both reviews collected data from some of the studies included in this review (Arroyo 2004; Beydoun 2005; Elger 2005; French 2003). Although both studies are in overall agreement with our current review, our current review provides novel information compared to the currently available reviews due to the meta-analysis conducted and the additional studies included.

The latter review by Ryvlin 2008 also discussed the long-term effectiveness of pregabalin after reviewing data collected from four open-label extension studies. Collectively, the data suggested that there was no loss of efficacy with the long-term use of pregabalin. For participants who entered long-term extension studies, 3.7% of people remained seizure-free during the last year of the respective studies. Likewise, a long-term observational study that followed 105 people (aged 16 to 81 years) over a one-year period revealed that 5.7% of people reported that they had been seizure-free for the previous four weeks when contacted at 12 months, and 17.1% of people reported that they had a 50% or greater reduction in seizure frequency over the 12-month period (Brandt 2009). Although all the studies included in our review were of short duration (treatment periods varied from 12 to 14 weeks for all outcomes), these observations suggest that there should be no decline in efficacy over longer time periods. However, this hypothesis remains to be investigated and demonstrated by randomised controlled trials of longer durations.

Another review specifically compared the efficacy of pregabalin and gabapentin by conducting a meta-analysis and by performing an indirect comparison method (Delahoy 2010). The review reported that pregabalin was more efficacious than gabapentin. Specifically, at the highest doses of both drugs, 600 mg pregabalin versus 1800 mg gabapentin, Delahoy 2010 reported an odds ratio of 2.52 (95% CI 1.21 to 5.27) for the outcome 50% or greater reduction in seizure frequency, in favour of pregabalin.

The findings of the review by Delahoy 2010 are in contrast to those of the randomised controlled trial conducted by French 2016 that we included in this review, and which directly compared pregabalin and gabapentin. French 2016 demonstrated no significant difference in efficacy between the two drugs for the outcome 50% or greater reduction in seizure frequency (RR 0.96, 95% CI 0.82 to 1.12). Furthermore, French 2016 reported that participants randomised to either of the two treatment groups (median dose 450 mg/d pregabalin and median dose 1500 mg/d gabapentin) experienced comparable percentage reductions in seizure frequency (58.7% and 57.4% median per cent reduction in seizure frequency, respectively).

Notably, the review by Delahoy 2010 has the advantage of including data from multiple sources and, as a result, includes a larger sample size. This should, in theory, provide a more accurate estimate of the effect of the two drugs. The study by French 2016, however, has the benefit of being a direct comparison between the two drugs which provides more convincing evidence than an indirect comparison. Specifically, the randomised controlled trial would be expected to have a more even distribution of participant characteristics at baseline and would have used a more standardised approach to compare the two drugs than the review. More randomised controlled trials directly comparing the two drugs would be necessary to appropriately compare their effectiveness and to enable conclusive findings to be reached.

With regard to the adverse event profile of pregabalin, the two reviews discussed earlier both reported somnolence, dizziness, ataxia, and fatigue as the most commonly reported adverse effects (Hamandi 2006; Ryvlin 2008), in keeping with the findings in this review. Ryvlin 2008 also specified that most adverse effects were mild to moderate in severity. Additionally, Ryvlin 2008 observed a dose-response relationship in the reporting of adverse effects. Weight gain was also a common adverse effect, with Ryvlin 2008 reporting that 24% of participants experienced weight gain whilst receiving pregabalin. In actuality, weight gain was the most reported adverse effect for people who participated in the observational study by Brandt 2009, followed by tiredness and cognitive disturbances.

In another meta-analysis specifically focused on investigating the adverse event profile of pregabalin (Zaccara 2012), it was shown that vestibulo-cerebellar and central nervous system adverse events, including ataxia and somnolence, were more commonly reported when pregabalin was used in those with focal epilepsy than when used for its other clinical indications, including anxiety disorders and pain disorders. It was suggested that these adverse events may not necessarily be attributable to pregabalin, but could instead be associated with an individual's concomitant AEDs or could actually be a symptom of focal epilepsy itself.

In contrast, Ryvlin 2008 appeared to consider the adverse effects reported by people with epilepsy to be a true representation of

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the adverse event profile of pregabalin. Ryvlin 2008 suggested that tolerability amongst people with epilepsy can instead be improved by individualising the dose of pregabalin, namely by using a titration and dose adjustment protocol, to limit adverse effects. Specifically, Ryvlin 2008 recognised that 24% of participants withdrew from trials during the first week of treatment in studies that utilised a fixed dose of pregabalin. This was compared to a withdrawal rate of only 3% in studies that used an individualised flexible-dose regimen. In studies of short duration, adverse effects are the most common reason for treatment withdrawal. Although this trend was not recognised in our current review, this outcome, that is treatment withdrawal within a given time period, may be of interest for future review updates as it would be informative for clinical practice.

AUTHORS' CONCLUSIONS

Implications for practice

In the relatively short term (12 to 17 weeks), pregabalin, given in a twice- or three-times-daily regimen, can significantly reduce seizure frequency in adults and children with treatment-resistant focal epilepsy. A dose of 600 mg/d immediate-release pregabalin can also significantly increase seizure freedom rates amongst people with treatment-resistant focal epilepsy, but is associated with a significantly higher treatment withdrawal rate compared to placebo. Pregabalin was significantly associated with the following adverse effects: ataxia, dizziness, fatigue, somnolence, headache, nausea and weight gain. The evidence suggests that there is no significant difference in efficacy and harms between pregabalin and some of the other currently available antiepileptic drugs, namely gabapentin, levetiracetam, and lamotrigine.

Implications for research

To improve clinical decisions, further clinical trials are required in adults and children with drug-resistant focal epilepsy. These trials should:

- 1. compare the efficacy and tolerability of pregabalin with other adjunctive treatments;
- 2. be of long-term duration (at least 12 months);
- 3. assess seizure freedom rates, quality of life, and health economic outcomes;
- 4. establish cost-effectiveness and compare it with that of other antiepileptic drugs.

Further data regarding pregnancy outcomes are also needed, which will require the recruitment of women taking pregabalin to ongoing pregnancy registries.

ACKNOWLEDGEMENTS

This review update was supported by the National Institute for Health Research, via Cochrane Infrastructure funding to the Epilepsy Group. The views and opinions expressed therein are those of the author(s) and do not necessarily reflect those of the Systematic Reviews Programme, NIHR, NHS or the Department of Health and Social Care.

We acknowledge Dora Lozsadi for contributions made in the original review, and to Jennifer Pulman and Karla Hemming for contributions to previous updates.

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Antinew 2019	
Study characteristics	
Methods	Randomised, double-blind, placebo-controlled, parallel-group, multicentre trial (18 countries: Bel- gium, Bulgaria, Czech Republic, France, Greece, Hungary, Israel, Italy, Malaysia, Philippines, Poland, Ro- mania, Serbia, Singapore, Korea, Turkey, Ukraine, and USA).
	3 treatment arms: 1 PBO, 1 PGB 2.5 mg/kg/d, and 1 PGB 10 mg/kg/d.

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ntinew 2019 (Continued)	
	Participants randomised to 1 of 3 treatment arms (capsules or liquid oral solution). Maximum dose was 150 mg/d for pregabalin 2.5 mg/kg/d, and 600 mg/d for pregabalin 10/mg/d.
	Duration of screening and baseline period: 8 weeks; 12-week double-blind treatment phase (2-week dose escalation and 10-week fixed dose) and 1-week taper phase.
Participants	Participants aged 4 to 16 years (mean 10.3 years), of either sex (57.4% male), all with treatment-resis- tant focal epilepsy. Participants were on 1 to 3 baseline AEDs.
	372 people screened, 295 participants randomised: 94 participants to PBO; 104 participants to 2.5 mg/ kg/d PGB; and 97 participants to 10 mg/kg/d PGB.
Interventions	Group 1: Placebo (no dose escalation)
	Group 2: PGB 2.5 mg/kg daily (no dose escalation)
	Group 3: PGB 10 mg/kg daily (2.5 mg/kg/d for week 1, 5 mg/kg/d for week 2).
Outcomes	Primary outcome: reduction in seizure frequency (50% responder status)
	Secondary outcomes: seizure freedom for maintenance phase (last 28-day seizure-free rates); adverse effects.
Notes	One participant randomised to the PBO group was excluded from ITT analysis because lost to fol- low-up.
	No information provided on methods of randomisation, concealment, or blinding.

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Participants randomised using 1:1:1 ratio. No further information given.
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	No information provided.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.
Selective reporting (re- porting bias)	Low risk	All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.
Other bias	High risk	Clinical Study was funded by Pfizer Inc.

Arroyo 2004

Study characteristics

Pregabalin add-on for drug-resistant focal epilepsy (Review)

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Arroyo 2004 (Continued)			
Methods	Randomised, double-blind, PBO-controlled, parallel, multicentre trial (45 in Europe, Australia, and South Africa). 3 treatment arms: 1 PBO, 2 PGB.		
	Participants randomised in blocks of 6, each allocated unique ID number. All participants received 2 capsules 3 times a day, but 2 capsule sizes were used (no further information available).		
	Duration of baseline period: 8 weeks. 12-week treatment period included 4- to 8-day titration period.		
Participants	Adults aged 17 to 73 years (mean 37 years), 50.5% male, all with drug-resistant focal epilepsy. Partici- pants were on 1 to 4 baseline AEDs.		
	344 people screened, 288 participants randomised: 97 participants to PBO (mean baseline 28-day seizure frequency: 23.5); 99 participants to 50 mg/d PGB 3 times a day (mean baseline 28-day seizure frequency: 26.2); and 92 participants to 200 mg/d PGB 3 times a day (mean baseline 28-day seizure frequency: 19.3).		
Interventions	Group 1: Placebo Group 2: PGB 50 mg 3 times a day (150 mg/d; 4-day titration phase) Group 3: PGB 200 mg 3 times a day (600 mg/d; 8-day titration phase)		
Outcomes	Primary outcome: reduction in seizure frequency compared to baseline (response ratio)		
	Secondary outcomes: responder rate, seizure freedom, change in seizure frequency, adverse effects		
Notes	Study used capsules of 2 sizes, containing 25 mg PGB or PBO (size 1# = small capsules) and 100 mg PGB or PBO (size 4# = large capsules). It is stated that participants received 2 capsules 3 times a day. 1 par- ticipant excluded from ITT in PBO arm, as failed to take study drugs.		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Random computer-generated code used stratified by centre using block size of 6.
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding (performance bias and detection bias) All outcomes	Low risk	Medication presented in identical capsules.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis performed.
Selective reporting (re- porting bias)	Low risk	All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.
Other bias	High risk	As an EEG was not required to confirm the above, some of the 18 participants included who were stated as having "generalised seizures", rather than sec- ondary generalised, may have had primary generalised epilepsy. Study was sponsored by Pfizer Inc.



Baulac 2010

Study characteristics		
Methods	Randomised, double-blind, PBO- and active-drug-controlled, parallel, multicentre trial (97 in Europe, Canada, and Australia).	
	3 treatment arms: 1 PBO, 1 PGB, and 1 LTG.	
	Participants randomised to 1 of 3 treatment arms (no further information available).	
	Duration of baseline period: 6 weeks. 17-week treatment period with 2 phases in addition to a titration phase (1 week of titration for PGB and 5 weeks of titration for LTG). (Phase I: 11 weeks treatment includ- ing 1 week titration for PGB and 5 weeks titration for LTG; Phase II: 6 weeks treatment).	
Participants	Adults aged 16 to 82 years (mean 39.4 years), 48.5% male, all with treatment-resistant focal epilepsy confirmed by history and recent EEG. Participants were on 1 to 3 baseline AEDs.	
	546 people screened, 434 participants randomised: 141 participants to PBO (mean baseline 28-day seizure frequency: 16.38); 152 participants to 150 mg to 300 mg PGB twice daily (mean baseline 28-day seizure frequency: 21.32); and 141 participants to 150 mg to 300 mg LTG twice daily (mean baseline 28-day day seizure frequency: 21.80).	
Interventions	Group 1: Placebo	
	Group 2: PGB 150 mg to 300 mg twice daily (300 to 600 mg/d; 1-week titration phase)	
	Group 3: LTG 150 mg to 300 mg twice daily (300 to 600 mg/d; 5-week titration phase)	
Outcomes	Primary outcome: change in seizure frequency compared to baseline (response ratio)	
	Secondary outcomes: responder rate, seizure freedom, adverse effects	
Notes	One participant randomised to the PBO group failed to take > 1 dose of medication and was therefore excluded from ITT analysis. No information provided on methods of randomisation, concealment, or blinding.	
	Study was sponsored by Pfizer Inc.	
Risk of bias		

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	No details of method of randomisation.
Allocation concealment (selection bias)	Unclear risk	No details of allocation concealment.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Same number of capsules administered per study day per group. No further details provided.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.
Selective reporting (re- porting bias)	Low risk	All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.
Other bias	High risk	Study was sponsored by Pfizer Inc.

Pregabalin add-on for drug-resistant focal epilepsy (Review)

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Beydoun 2005

Study characteristics			
Methods	Randomised, double-b	olind, PBO-controlled, parallel, multicentre trial (43 in USA and Canada).	
	3 treatment arms: 1 PB	30, 2 PGB.	
		ed in blocks of 6, each allocated unique ID number. All participants received 3- blinded capsules (no further information available).	
	Duration of baseline pe	eriod: 8 weeks. 12-week treatment period with 1-week titration period.	
Participants		ears (mean 39.1 years), 50.2% male, all with treatment-resistant focal epilepsy nd recent EEG. Participants were on 1 to 4 baseline AEDs.	
	378 people screened, 313 participants randomised: 98 participants to PBO (mean baseline 28-day seizure frequency: 25.1); 104 participants to 300 mg PGB twice daily (mean baseline 28-day seizure frequency: 21.5); and 111 participants to 200 mg PGB 3 times a day (mean baseline 28-day seizure frequency: 21.3).		
Interventions		wice daily (600 mg/d; 1-week titration phase) 8 times a day (600 mg/d; 1-week titration phase)	
Outcomes	Primary outcome: reduction in seizure frequency compared to baseline (response ratio)		
	Secondary outcomes:	responder rate, median percentage change in seizure frequency	
Notes		sed to the 300 mg twice-daily group failed to take tablets and was therefore ex- sis. Blinding was broken with 1 participant in the PBO arm when she became	
	Study was sponsored b	by Pfizer Inc.	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Participants randomised in blocks of 6 and allocated unique ID number.	
Allocation concealment (selection bias)	Unclear risk	No details provided.	
Blinding (performance bias and detection bias) All outcomes	Low risk	All participants received identical capsules.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.	

 Selective reporting (reporting (reporting bias)
 Low risk
 All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.

 Other bias
 High risk
 Study was sponsored by Pfizer Inc.

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Elger 2005

Study characteristics			
Methods	Randomised, double-b	lind, PBO-controlled, parallel, multicentre trial (53 in Europe and Canada).	
		30, 2 PGB. Participants randomised in blocks of 5, each allocated unique ID num- cked control group using identical capsules (no further information available).	
	Duration of baseline pe	eriod: 6 weeks. 12-week treatment period.	
Participants		ears (mean 40.5 years), 49.9% male, all with treatment-resistant focal epilepsy and family history as well as recent EEG.	
	ipants to PBO (median daily fixed dose (media	to 5 baseline AEDs. 400 people screened, 341 participants randomised: 73 partic- baseline 28-day seizure frequency: 8.7); 137 participants to 300 mg PGB twice- an baseline 28-day seizure frequency: 10); and 131 participants to PGB flexible ne 28-day seizure frequency: 9.33).	
Interventions	Group 1: Placebo Group 2: 300 mg PGB twice-daily fixed dose (600 mg/d) Group 3: 75 mg to 300 mg PGB twice-daily flexible titration at physician's discretion (150 to 600 mg/d)		
Outcomes	Primary outcome: reduction in seizure frequency compared to baseline (response ratio)		
		responder rate, median percentage change in seizure frequency and reduction of ing the study, adverse effects	
Notes	In PGB titration and PBO groups, participants were included with seizure frequency of over 120 a day Documenting seizures at this frequency is difficult and may be unreliable. Medium length of follow-unot reported.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Participants randomised using a 1:2:2 ratio and block sizes of 5.	
Allocation concealment (selection bias)	Unclear risk	No information provided.	
Blinding (performance bias and detection bias) All outcomes	Low risk	Study medication presented in identical capsules.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.	
Selective reporting (re-	Low risk	All outcomes stated in methods section of paper were reported in the results.	

 porting bias)
 However, there was no protocol available to check a priori outcomes.

 Other bias
 High risk
 Study was sponsored by Pfizer Inc.

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Study characteristics			
Methods	Randomised, double-b	lind, PBO-controlled, parallel, multicentre trial (71 in the USA and 5 in Canada).	
		O, 4 PGB. Participants randomised in blocks of 5, each allocated unique ID numed (no further information available).	
	Duration of baseline pe	eriod: 8 weeks. There was no titration; 12-week treatment period.	
Participants	Participants 12 years and above (range 12 to 75 years, mean 38.4 years), 48.1% male, all with treat- ment-resistant focal epilepsy. Participants were on 1 to 4 baseline AEDs.		
	586 people screened, 455 participants randomised: 100 participants to PBO (mean baseline seizure fre quency: 22.3); 88 participants to 50 mg PGB (mean baseline seizure frequency: 27.4); 88 participants to 150 mg PGB (mean baseline 28-day seizure frequency: 23.1); 90 participants to 300 mg PGB (mean base line 28-day seizure frequency: 19.1); and 89 participants to 600 mg PGB (mean baseline 28-day seizure frequency: 18.6).		
Interventions	Group 1: Placebo Group 2: 50 mg/d PGB Group 3: 150 mg/d PGE Group 4: 300 mg/d PGE Group 5: 600 mg/d PGE	3 (twice daily) 3 (twice daily)	
Outcomes	Primary outcome: redu	iction in seizure frequency compared to baseline (response ratio)	
	Secondary outcomes: responder rate, pairwise comparisons with PBO, adverse effects		
Notes	Blinding broken for interim analysis (data obtained were only known to committee who were not in- volved in further running of study) and for 1 participant who developed visual field defect. Two partici- pants were excluded from ITT analysis (1 withdrew consent, 1 had AEDs changed during baseline peri- od). Seizure frequency and responder rate were calculated from data collected from seizure diaries an mean calculated over a 4-week period.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Used a computer-generated randomised schedule using block sizes of 5.	
Allocation concealment (selection bias)	Unclear risk	No information provided.	
Blinding (performance bias and detection bias) All outcomes	Low risk	Study medication presented in identical capsules.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.	
Selective reporting (re- porting bias)	Low risk	All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.	
Other bias	High risk	Possibility of the inclusion of individuals with primary generalised epilepsy.	
		Study was sponsored by Pfizer Inc.	

Pregabalin add-on for drug-resistant focal epilepsy (Review)



French 2014

Study characteristics			
Methods		olind, placebo-controlled, parallel, multicentre trial (18 countries) assessing the y of controlled-release PGB.	
	3 treatment arms: 1 PB	30, 2 PGB	
	Randomised 1:1:1 to Pesation system.	GB 165 mg/d or PGB 330 mg/d or placebo using a computer-generated randomi-	
		eriod: 8 weeks. 14-week double-blind treatment period with 2-week double-blind on phase); 1-week taper.	
Participants	Adults aged 18 to 75 years, 47.7% male, all with focal epilepsy. Participants were on 1 to 3 baseline AEDs.		
	400 people screened, 325 participants randomised: 110 participants to placebo (mean baseline 28- day seizure frequency: 17.8); 101 participants to PGB 165 mg (mean baseline 28-day seizure frequency: 13.0); 114 participants to PGB 330 mg (mean baseline 28-day seizure frequency: 17.0).		
Interventions	Group 1: Placebo		
	Group 2: PGB 165 mg/d controlled release Group 3: PGB 330 mg/d controlled release		
Outcomes	Primary outcomes: reduction in seizure frequency compared to baseline (response ratio)		
	Secondary outcomes: responder rate, adverse effects		
Notes	Clinical trials: NCT01262677		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated randomisation system.	
Allocation concealment (selection bias)	Unclear risk	Allocate participants to each of the 3 treatment groups in a 1:1:1 manner. No further details provided.	
Blinding (performance bias and detection bias) All outcomes	Unclear risk	No details regarding blinding of participants and personnel provided.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT efficacy analysis performed.	
Selective reporting (re-	Low risk	All outcomes stated in methods section of paper were reported in the results.	
porting bias)		There was no protocol available to check a priori outcomes.	
Other bias	High risk	Study sponsored by Pfizer Inc.	

French 2016

lestern Europe, Asia, on system. uble-blind dose esca-
uble-blind dose esca-
lequately controlled
ng median dose (mean n dose (mean baseline
se) tion phase)
res and 75% or more
-free rates), adverse
1

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Computer-generated randomisation system.
Allocation concealment (selection bias)	Unclear risk	Allocate participants to each of the 2 treatment groups in a 1:1 manner. No fur- ther details provided.
Blinding (performance bias and detection bias) All outcomes	Low risk	Medication presented in identical tablets. Identical analysis of results.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT efficacy analysis performed.
Selective reporting (re-	Low risk	All outcomes stated in methods section of paper were reported in the results.
porting bias)		There was no protocol available to check a priori outcomes.
Other bias	High risk	Study was sponsored by Pfizer Inc.

Lee 2009

Study characteristics

Pregabalin add-on for drug-resistant focal epilepsy (Review)

ee 2009 (Continued)			
Methods	Randomised, double-blind, PBO-controlled, parallel, multicentre trial (9 in Korea).		
	2 treatment arms: 1 PB	30, 1 PGB	
	Participants randomise	ed to 1 of 2 treatment arms (no further information available).	
	Duration of baseline pe	eriod: 6 weeks. 12-week treatment period (no further details provided).	
Participants		nd above (mean 34.2 years), 48.3% male, all with treatment-resistant focal were on 1 to 3 baseline AEDs.	
	209 people screened, 178 participants randomised: 59 participants to PBO (mean baseline 28-day seizure frequency: 13.2) and 119 participants to 150 mg to 600 mg PGB (mean baseline 28-day seizure frequency 13.2).		
Interventions	Group 1: Placebo		
	Group 2: 75 mg to 300 i	mg PGB twice daily (150 to 600 mg/d)	
Outcomes	Primary outcome: change in seizure frequency (response ratio)		
	Secondary outcomes: verse effects	responder rate, seizure freedom, anxiety/depression, sleep, quality of life, ad-	
Notes	All randomised participants included in ITT analysis. No information provided on methods of random sation, concealment, or blinding.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Unclear risk	Participants randomised using 2:1 ratio. No further information given.	
Allocation concealment (selection bias)	Unclear risk	No information provided.	
Blinding (performance bias and detection bias) All outcomes	Unclear risk	No information provided.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT analysis employed.	
Selective reporting (re- porting bias)	Low risk	All outcomes stated in methods section of paper were reported in the results. However, there was no protocol available to check a priori outcomes.	

Mann 2020

Other bias

Study characteristics

Methods

Randomised, double-blind, placebo-controlled, parallel-group, multicentre trial (22 countries: Belarus, Belgium, Bulgaria, China, France, Germany, Greece, Hungary, Israel, Lebanon, Malaysia, Philippines, Re-

Study was sponsored by Pfizer Inc.

Pregabalin add-on for drug-resistant focal epilepsy (Review)

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High risk

3 treatment arms: 1 PGB 7 mg/kg/d, PGB 14 mg/kg/d, and 1 PBO (pregabalin or matching placebo was administered as a 20-mg/mL oral solution in 3 equally divided doses per day). Participants randomised to 1 of 3 treatment arms. 48 to 72 hours baseline phase; 14-day double-blind treatment phase (5-day dose escalation, 99-day finated dose including V-EEG monitoring of 48-72 hours over final 3 days) and a 7-day double-blind taper phase. Participants Children aged 1 month to 4 years (mean 28.2 months), of either sex (59% male), all with treatment-resistant focal epilepsy. Participants randomised: 71 participants to 7 mg/kg/d PGB; 34 participants to 14 mg/kg/d PGB; 34 participants of 7 mg/kg/d PGB; 34 participants to 14 mg/kg/d PGB; 34 participants randomised: 71 participants to 7 ug/kg/d PGB; 34 participants to 14 mg/kg/d PGB; 34 participants randomised: 71 participants to 7 ug/kg/d PGB; 34 participants to 14 mg/kg/d PGB; 34 participants randomised: 71 participants to 7 ug/kg/d PGB; 34 participants to 14 mg/kg/d PGB; 34 participants randomised: 71 participants to 7 ug/kg/d PGB; 34 participants of age; stratum 3, 1 < Year of age; st	Mann 2020 (Continued)	public of Korea, Romar USA).	nia, Russian Federation, Serbia, Spain, Taiwan, Thailand, Turkey, Ukraine, and
48 to 72 hours baseline phase; 14-day double-blind treatment phase (5-day dose escalation, 99-day fixed dose including V-EEG monitoring of 48-72 hours over final 3 days) and a 7-day double-blind taper phase. Participants Children aged 1 month to 4 years (mean 28.2 months), of either sex (59% male), all with treatment-re-sistant focal epilepsy. Participants were on 1 to 3 baseline AEDs. 231 people screened, 175 participants randomised: 71 participants to 7 mg/kg/d PGB; 34 participants to 14 mg/kg/d PGB; and 70 participants to PEO. Randomization was stratified by study site and participant age strata as follows: stratum 1, < 1 year of age; stratum 2, 1 to 2 years of age; stratum 3, > 2 years of age; There was no prespecified number to be enrolled per stratum. Interventions Group 1: Placebo Group 2: Pregabalin 7 mg/kg daily Group 3: Pregabalin 14 mg/kg daily Outcomes Primary outcome: reduction in seizure frequency (50% responder rate) Secondary outcome: adverse effects. Secondary outcomes: adverse effects. Notes ClinicalTrials.gov registration: NCT02072824. Risk of bias Support for judgement Bias Authors' judgement Support for judgement Allocation concealment (selection bias) Low risk Telerandomisation system according to the agreed randomisation code. Bilaiding (performance bias and detection bias) Low risk The double-blind fixed-dose treatment was administered by parentits/guaradins/j/caregiver(s) according to provided inst			
fixed dose including V-EEG monitoring of 48-72 hours over final 3 days) and a 7-day double-blind taper phase. Participants Children aged 1 month to 4 years (mean 28.2 months), of either sex (59% male), all with treatment-resistant focal epilepsy. Participants were on 1 to 3 baseline AEDs. 231 people screened, 175 participants randomised: 71 participants to 7 mg/kg/d PGB; 34 participants to 14 mg/kg/d PGB, and 70 participants to PBO. Randomization was stratified by study site and participant age stratatum 1, < 1 year of age; stratum 2, 1 to 2 years of age; stratum 3, > 2 years of age. There was no prespecified number to be enrolled per stratum. Interventions Group 1: Placebo Group 2: Pregabalin 7 mg/kg daily Group 3: Pregabalin 7 mg/kg daily Outcomes Primary outcome: reduction in seizure frequency (50% responder rate) Secondary outcome: adverse effects. Notes Notes ClinicalTrials.gov registration: NCT02072824. Risk of bias Authors' judgement Support for judgement Allocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants. Blinding (performance bias) Low risk Allocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants. Blinding (performance bias and betection bias) Low risk Allocated participants coercing to provided instructions from day 6 to day 15.		Participants randomise	ed to 1 of 3 treatment arms.
sistant focal epilepsy. Participants were on 1 to 3 baseline AEDs. 231 people screened, 175 participants to 7 mg/kg/d PGB; 34 participants to 14 mg/kg/d PGB; and 70 participants to PBO. Randomization was stratified by study site and participant age strata as follows: stratum 7, 1 to 2 years of age; stratum 3, > 2 years of age. There was no prespecified number to be enrolled per stratum. Interventions Group 1: Placebo Group 2: Pregabalin 7 mg/kg daily Outcomes Primary outcome: reduction in seizure frequency (50% responder rate) Secondary outcomes: adverse effects. Notes ClinicalTrials.gov registration: NCT02072824. Bias Authors' judgement Authors' judgement Support for judgement Allocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants. Blinding (performance bias) Low risk The double-bilind fixed-dose treatment was administered by parentis/jucardian(s)/caregiver(s) according to provided instructions from day 6 to day 15. Incording isolary Low risk Attrition rates reported. ITT efficacy analysis performed. All outcomes Low risk All outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		fixed dose including V-	
to 14 mg/kg/d FG8; and To participants to PB0. Randomization was stratified by study site and participant age strata as follows: stratum 1, < 1 year of age; stratum 2, 1 to 2 years of age; stratum 3, > 2 years of age;	Participants		
Group 2: Pregabalin 7 mg/kg daily Group 3: Pregabalin 14 mg/kg dailyOutcomesPrimary outcome: reduction in seizure frequency (50% responder rate) Secondary outcomes: adverse effects.NotesClinicalTrials.gov registration: NCT02072824. Risk of biasAuthors' judgement BiasAuthors' judgementRandom sequence generation (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Blinding (performance 		to 14 mg/kg/d PGB; an pant age strata as follo	d 70 participants to PBO. Randomization was stratified by study site and partici- ws: stratum 1, < 1 year of age; stratum 2, 1 to 2 years of age; stratum 3, > 2 years
Group 3: Pregabalin 14 mg/kg dailyOutcomesPrimary outcome: reduction in seizure frequency (50% responder rate) Secondary outcomes: adverse effects.NotesClinicalTrials.gov registration: NCT02072824.Risk of biasSupport for judgementBiasAuthors' judgementRandom sequence generation (selection bias)Low riskAllocation concealment (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Binding (performance 	Interventions	Group 1: Placebo	
OutcomesPrimary outcome: reduction in seizure frequency (50% responder rate) Secondary outcomes: adverse effects.NotesClinicalTrials.gov registration: NCT02072824.Risk of biasAuthors' judgementSupport for judgementBiasAuthors' judgementSupport for judgementRandom sequence generation (selection bias)Low riskTelerandomisation system according to the agreed randomisation code.Allocation concealment (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Blinding (performance bias and detection bias)Low riskThe double-blind fixed-dose treatment was administered by paren- t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		Group 2: Pregabalin 7 r	ng/kg daily
Secondary outcomes: adverse effects.NotesClinicalTrials.gov registration: NCT02072824.Risk of biasAuthors' judgementSupport for judgementBiasAuthors' judgementSupport for judgementRandom sequence generation (selection bias)Low riskTelerandomisation system according to the agreed randomisation code.Allocation concealment (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Blinding (performance bias and detection bias)Low riskThe double-blind fixed-dose treatment was administered by paren- t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		Group 3: Pregabalin 14	mg/kg daily
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Risk of biasBiasAuthors' judgementSupport for judgementRandom sequence generation (selection bias)Low riskTelerandomisation system according to the agreed randomisation code.Allocation concealment (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Blinding (performance bias and detection bias)Low riskThe double-blind fixed-dose treatment was administered by paren- t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		Secondary outcomes:	adverse effects.
BiasAuthors' judgementSupport for judgementRandom sequence generation (selection bias)Low riskTelerandomisation system according to the agreed randomisation code.Allocation concealment (selection bias)Low riskAllocated participants to each of the 3 treatment groups in a 2:1:2 manner. The dose was based on the age and weight of the participants.Blinding (performance bias and detection bias)Low riskThe double-blind fixed-dose treatment was administered by paren- t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.	Notes	ClinicalTrials.gov regis	tration: NCT02072824.
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(selection bias)dose was based on the age and weight of the participants.Blinding (performance bias and detection bias) All outcomesLow riskThe double-blind fixed-dose treatment was administered by paren- t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		Low risk	Telerandomisation system according to the agreed randomisation code.
bias and detection bias) All outcomest(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to day 15.Incomplete outcome data (attrition bias) All outcomesLow riskAttrition rates reported. ITT efficacy analysis performed.Selective reporting (re- porting bias)Low riskAll outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.		Low risk	
(attrition bias) All outcomes Selective reporting (re- porting bias) Low risk All outcomes stated in methods section of paper were reported in the results. There was no protocol available to check a priori outcomes.	bias and detection bias)	Low risk	t(s)/guardian(s)/caregiver(s) according to provided instructions from day 6 to
porting bias) There was no protocol available to check a priori outcomes.	(attrition bias)	Low risk	Attrition rates reported. ITT efficacy analysis performed.
There was no protocol available to check a priori outcomes.		Low risk	All outcomes stated in methods section of paper were reported in the results.
Other bias High risk Study sponsored by Pfizer Inc.	porting bias)		There was no protocol available to check a priori outcomes.
	Other bias	High risk	Study sponsored by Pfizer Inc.

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Zaccara 2014

Study characteristics			
Methods	Randomised, double-b Europe, South and Cer	lind, flexible-dose, parallel, multicentre trial (71 centres in Western and Eastern htral America, Asia).	
	2 treatment arms: PGB	and LEV	
	Randomised 1:1 to eith	ner PGB or LEV using a computer-generated randomisation system.	
	Duration of baseline pl double-blind maintena	hase: 6 weeks; 4-week double-blind dose escalation (titration phase); 12-week ance phase.	
Participants		ears (mean 37 years), all with drug-resistant focal epilepsy (inadequately con- but no more than 5 AEDs). Participants were on 1 to 2 baseline AEDs.	
		ned, 509 participants were randomised: 254 participants to PGB (mean baseline icy: 16.2) and 255 participants to LEV (mean baseline 28-day seizure frequency:	
Interventions		ily (150, 300, 450, and 600 mg/d) ly (1000, 2000, and 3000 mg/d)	
Outcomes	Primary outcome: reduction in seizure frequency (50% or more reduction of seizures)		
	Secondary outcomes: seizure freedom for maintenance phase, adverse effects		
Notes	Clinical trials: NCT00537238		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Computer-generated randomisation system.	
Allocation concealment (selection bias)	Unclear risk	Allocated participants to each of the 2 treatment group in a 1:1 ratio. No fur- ther details provided.	
Blinding (performance bias and detection bias)	Low risk	Details are not provided. However, it is likely that blinding of participants and personnel was maintained due to the methods used.	
All outcomes		Medication presented in identical tablets. Identical analysis of results.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition rates reported. ITT efficacy analysis performed.	
Selective reporting (re-	Low risk	All outcomes stated in methods section of paper were reported in the results.	
porting bias)		No protocol available to check a priori outcomes.	
Other bias	High risk	Study was sponsored by Pfizer Inc.	

AED: antiepileptic drug; EEG: electroencephalogram; GPN: gabapentin; GTCS: generalised tonic-clonic seizures; ITT: intention-to-treat; LEV: levetiracetam; LTG: lamotrigine; PBO: placebo; PGB: pregabalin; V-EEG: video-electroencephalogram

Characteristics of excluded studies [ordered by study ID]

Pregabalin add-on for drug-resistant focal epilepsy (Review)

Study	Reason for exclusion	
Aicua-Rapun 2020	The study analysed other types of outcome measures.	
Hu 2020	Not an RCT.	
Morano 2019	Not an RCT.	
Moseley 2019	Not an RCT.	
Taghdiri 2015	This previously ongoing study, IRCT2012091210508N4, was published and excluded from the cur- rent review as the participant sample (including participants with a diagnosis of primary gener- alised epilepsy) did not meet our inclusion criteria.	

Characteristics of studies awaiting classification [ordered by study ID]

Russi 2006	
Methods	Randomised observational controlled study. 4 treatment arms: 1) LEV fast-rate, 2) LEV slow-rate, 3) PGB fast-rate, 4) PGB slow-rate dosage
Participants	128 participants with refractory focal epilepsy (32 in each treatment arm)
Interventions	Group 1: starting dose of 1000 mg twice daily LEV fast rate with weekly increments of 500 mg
	Group 2: starting dose of 500 mg twice daily LEV slow rate with weekly increments of 250 mg
	Group 3: starting dose of 300 mg twice daily PGB fast rate with weekly increments of 150 mg
	Group 4: starting dose of 150 mg twice daily PGB slow rate with weekly increments of 75 mg
Outcomes	Rate of withdrawals and continuation to maximum dose
	Incidence of adverse effects
Notes	Study reported in abstract form only. Further details of study are unavailable.

Tata 2007

Methods	Randomised cross-over trial consisting of 2 treatment arms: 1) PGB, 2) LEV. Participants ran- domised to groups using 1:1 ratio. Study was open-label. Long-term study duration of minimum 6 months.	
Participants	28 adults aged 19 to 62 years, 54% male. Participants currently taking different AED without main- taining good seizure control, stabilised to therapeutic association of valproate and lamotrigine.	
Interventions	Group 1: starting dose of 150 mg to target dose of 600 mg PGB Group 2: starting dose of 1000 mg to target dose of 3000 mg LEV	
Outcomes	Seizure freedom Seizure reduction	
	Withdrawals	

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Tata 2007 (Continued)

Adverse effects

Notes

Study reported in abstract only. Further details of study are unavailable.

AED: antiepileptic drug; LEV: levetiracetam; PGB: pregabalin

DATA AND ANALYSES

Comparison 1. Pregabalin versus placebo

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.1 50% or greater reduction in seizure frequency - ITT	9	2663	Risk Ratio (IV, Random, 95% CI)	1.95 [1.40, 2.72]
1.2 50% or greater reduction in seizure frequency - best-case analysis	9	2663	Risk Ratio (IV, Random, 95% CI)	2.91 [1.92, 4.42]
1.3 50% or greater reduction in seizure frequency - worst-case analysis	9	2663	Risk Ratio (IV, Random, 95% CI)	1.10 [0.92, 1.31]
1.4 Seizure freedom	4	1125	Risk Ratio (M-H, Fixed, 95% CI)	3.94 [1.50, 10.37]
1.5 Treatment withdrawal for any reason	9	2663	Risk Ratio (M-H, Fixed, 95% CI)	1.33 [1.10, 1.60]
1.6 Treatment withdrawal due to adverse effects	9	2663	Risk Ratio (M-H, Fixed, 95% CI)	2.60 [1.86, 3.64]
1.7 Ataxia	6	1868	Risk Ratio (M-H, Fixed, 99% CI)	3.90 [2.05, 7.42]
1.8 Dizziness	7	2193	Risk Ratio (M-H, Fixed, 99% CI)	3.15 [2.23, 4.44]
1.9 Fatigue	8	2488	Risk Ratio (M-H, Fixed, 99% CI)	1.35 [0.94, 1.93]
1.10 Headache	6	1850	Risk Ratio (M-H, Fixed, 99% CI)	0.65 [0.45, 0.94]
1.11 Nausea	4	1267	Risk Ratio (M-H, Fixed, 99% CI)	1.20 [0.56, 2.58]
1.12 Somnolence	9	2663	Risk Ratio (M-H, Fixed, 99% CI)	2.05 [1.49, 2.81]
1.13 Weight gain	8	2488	Risk Ratio (M-H, Fixed, 99% CI)	4.35 [2.34, 8.11]



Analysis 1.1. Comparison 1: Pregabalin versus placebo, Outcome 1: 50% or greater reduction in seizure frequency - ITT

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Antinew 2019	69	201	21	94	12.1%	1.54 [1.01 , 2.34]	
Arroyo 2004	54	191	6	97	7.9%	4.57 [2.04 , 10.25]	_
Baulac 2010	54	152	30	141	12.5%	1.67 [1.14 , 2.45]	
Beydoun 2005	98	215	9	98	9.6%	4.96 [2.62 , 9.41]	
Elger 2005	103	268	8	73	9.3%	3.51 [1.79 , 6.86]	_
French 2003	121	355	14	100	11.1%	2.43 [1.47 , 4.04]	
French 2014	88	215	39	110	13.4%	1.15 [0.86 , 1.56]	
Lee 2009	55	119	19	59	12.1%	1.44 [0.94 , 2.18]	
Mann 2020	33	105	22	70	11.8%	1.00 [0.64 , 1.56]	-+-
Total (95% CI)		1821		842	100.0%	1.95 [1.40 , 2.72]	
Total events:	675		168				
Heterogeneity: Tau ² = 0).19; Chi ² = 3	5.47, df =	8 (P < 0.00	01); I ² = 7	7%		-++++++++++++++++++++++++++++++++++++
Test for overall effect: 2	Z = 3.97 (P <	0.0001)					Favours placebo Favours pregabalir

Test for subgroup differences: Not applicable

Analysis 1.2. Comparison 1: Pregabalin versus placebo, Outcome 2: 50% or greater reduction in seizure frequency - best-case analysis

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Antinew 2019	95	201	21	94	11.8%	2.12 [1.41 , 3.17]	+
Arroyo 2004	88	191	6	97	9.0%	7.45 [3.38 , 16.41]	
Baulac 2010	100	152	30	141	12.2%	3.09 [2.21 , 4.33]	-
Beydoun 2005	157	215	9	98	10.2%	7.95 [4.24 , 14.90]	
Elger 2005	191	268	8	73	9.9%	6.50 [3.37 , 12.56]	
French 2003	185	355	14	100	11.1%	3.72 [2.27 , 6.11]	
French 2014	112	215	39	110	12.5%	1.47 [1.11 , 1.95]	-
Lee 2009	65	119	19	59	11.8%	1.70 [1.13 , 2.54]	
Mann 2020	36	105	22	70	11.6%	1.09 [0.71 , 1.69]	+
Total (95% CI)		1821		842	100.0%	2.91 [1.92 , 4.42]	
Total events:	1029		168				•
Heterogeneity: Tau ² = 0).34; Chi ² = 6	2.88, df =	8 (P < 0.00	001); I ² =	87%		0.01 0.1 1 10 100
Test for overall effect: 2	Z = 5.03 (P <	0.00001)					Favours placebo Favours pregabalin

Test for subgroup differences: Not applicable



Analysis 1.3. Comparison 1: Pregabalin versus placebo, Outcome 3: 50% or greater reduction in seizure frequency - worst-case analysis

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Antinew 2019	69	201	31	94	11.4%	1.04 [0.74 , 1.47]	-
Arroyo 2004	54	191	19	97	8.5%	1.44 [0.91 , 2.29]	-
Baulac 2010	54	152	65	141	13.4%	0.77 [0.58 , 1.02]	-
Beydoun 2005	98	215	26	98	10.9%	1.72 [1.20 , 2.46]	-
Elger 2005	103	268	25	73	11.2%	1.12 [0.79 , 1.60]	+
French 2003	121	355	27	100	11.1%	1.26 [0.89 , 1.80]	-
French 2014	88	215	51	110	14.1%	0.88 [0.68 , 1.14]	4
Lee 2009	55	119	21	59	10.0%	1.30 [0.88 , 1.93]	
Mann 2020	33	105	25	70	9.3%	0.88 [0.58 , 1.34]	-
Total (95% CI)		1821		842	100.0%	1.10 [0.92 , 1.31]	
Total events:	675		290				•
Heterogeneity: Tau ² = 0	.04; Chi ² = 1	8.39, df =	8 (P = 0.02); I ² = 56%	6		0.01 0.1 1 10 100
Test for overall effect: Z	Z = 1.06 (P =	0.29)					Favours placebo Favours pregabalin
Test for subgroup differ	ences: Not a	pplicable					

Analysis 1.4. Comparison 1: Pregabalin versus placebo, Outcome 4: Seizure freedom

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Beydoun 2005	17	215	0	98	11.5%	16.04 [0.97 , 264.07]	
Baulac 2010	6	152	1	141	17.4%	5.57 [0.68 , 45.66]	
Elger 2005	8	268	1	73	26.3%	2.18 [0.28 , 17.14]	
Lee 2009	5	119	2	59	44.8%	1.24 [0.25 , 6.20]	_
Total (95% CI)		754		371	100.0%	3.94 [1.50 , 10.37]	
Total events:	36		4				-
Heterogeneity: Chi ² = 3.	.37, df = 3 (I	P = 0.34);]	[2 = 11%				0.01 0.1 1 10 100
Test for overall effect: Z	= 2.78 (P =	0.006)					Favours placebo Favours pregabalin
Test for subgroup differe	ences: Not a	pplicable					

Librarv

Analysis 1.5. Comparison 1: Pregabalin versus placebo, Outcome 5: Treatment withdrawal for any reason

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Antinew 2019	26	201	10	94	8.5%	1.22 [0.61 , 2.42]	
Arroyo 2004	34	191	13	97	10.8%	1.33 [0.74 , 2.40]	_ _
Baulac 2010	46	152	35	141	22.7%	1.22 [0.84 , 1.77]	
Beydoun 2005	59	215	17	98	14.6%	1.58 [0.98 , 2.57]	
Elger 2005	88	268	17	73	16.7%	1.41 [0.90 , 2.21]	+ - -
French 2003	64	355	13	100	12.7%	1.39 [0.80 , 2.41]	
French 2014	24	215	12	110	9.9%	1.02 [0.53 , 1.97]	_ _
Lee 2009	10	119	2	59	1.7%	2.48 [0.56 , 10.95]	_ _
Mann 2020	3	105	3	70	2.3%	0.67 [0.14 , 3.21]	
Total (95% CI)		1821		842	100.0%	1.33 [1.10 , 1.60]	
Total events:	354		122				•
Heterogeneity: Chi ² = 2	2.88, df = 8 (I	P = 0.94); 1	$1^2 = 0\%$			0.01	0.1 1 10 100
Test for overall effect: 2	Z = 2.91 (P =	0.004)				Favou	rs pregabalin Favours placebo
Test for subgroup differ	rences: Not a	pplicable					

Analysis 1.6. Comparison 1: Pregabalin versus placebo, Outcome 6: Treatment withdrawal due to adverse effects

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
Antinew 2019	5	201	0	94	1.3%	5.17 [0.29 , 92.60]	
Arroyo 2004	27	191	6	97	15.7%	2.29 [0.98 , 5.35]	
Baulac 2010	24	152	10	141	20.5%	2.23 [1.10 , 4.49]	
Beydoun 2005	48	215	7	98	19.0%	3.13 [1.47 , 6.66]	
Elger 2005	61	268	5	73	15.5%	3.32 [1.39 , 7.97]	
French 2003	41	355	5	100	15.4%	2.31 [0.94 , 5.69]	
French 2014	11	215	3	110	7.8%	1.88 [0.53 , 6.59]	
Lee 2009	7	119	0	59	1.3%	7.50 [0.44 , 129.13]	
Mann 2020	0	105	1	70	3.5%	0.22 [0.01 , 5.40]	,
Total (95% CI)		1821		842	100.0%	2.60 [1.86 , 3.64	1	
Total events:	224		37				•	
Heterogeneity: Chi ² = 4	4.16, df = 8 (I	P = 0.84);]	$I^2 = 0\%$				0.01 0.1 1 10	100
Test for overall effect:	Z = 5.56 (P <	0.00001)					Favours pregabalin Favours p	
Test for subgroup diffe							1 0 1	

Test for subgroup differences: Not applicable



	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Arroyo 2004	18	191	3	97	17.1%	3.05 [0.63 , 14.70]	· _ • _ • _ · · · · · · · · · · · · · ·
Baulac 2010	13	152	1	141	4.4%	12.06 [0.85 , 171.72]	I→
Beydoun 2005	48	215	6	98	35.3%	3.65 [1.25 , 10.63]	I
Elger 2005	41	268	3	73	20.2%	3.72 [0.83 , 16.73]	∣ – –
French 2003	34	355	3	100	20.1%	3.19 [0.70 , 14.65]	Ⅰ
Lee 2009	5	119	0	59	2.9%	5.50 [0.13 , 241.69]	l► →
Total (99% CI)		1300		568	100.0%	3.90 [2.05 , 7.42]	
Total events:	159		16				•
Heterogeneity: Chi ² = 1.	.56, df = 5 (I	P = 0.91);]	$1^2 = 0\%$				0.01 0.1 1 10 100
Test for overall effect: Z	z = 5.44 (P <	0.00001)					Favours pregabalin Favours placebo
Test for subgroup different	ences: Not a	pplicable					

Analysis 1.7. Comparison 1: Pregabalin versus placebo, Outcome 7: Ataxia

Analysis 1.8. Comparison 1: Pregabalin versus placebo, Outcome 8: Dizziness

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Arroyo 2004	43	191	8	97	13.7%	2.73 [1.07 , 6.98]	
Baulac 2010	38	152	13	141	17.4%	2.71 [1.25 , 5.86]	
Beydoun 2005	88	215	14	98	24.8%	2.87 [1.46 , 5.61]	
Elger 2005	91	268	6	73	12.2%	4.13 [1.47 , 11.58]	
French 2003	88	355	9	100	18.1%	2.75 [1.17 , 6.46]	
French 2014	22	215	2	110	3.4%	5.63 [0.86 , 36.82]	
Lee 2009	46	119	6	59	10.4%	3.80 [1.34 , 10.76]	_
Total (99% CI)		1515		678	100.0%	3.15 [2.23 , 4.44]	
Total events:	416		58				•
Heterogeneity: Chi ² = 2	.01, df = 6 (I	P = 0.92); 1	$2^2 = 0\%$			0	0.01 0.1 1 10 100
Test for overall effect: Z	Z = 8.59 (P <	0.00001)				Fa	vours pregabalin Favours placebo

Test for subgroup differences: Not applicable



	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Antinew 2019	10	201	3	94	4.9%	1.56 [0.29 , 8.24]
Arroyo 2004	26	191	11	97	17.4%	1.20 [0.50 , 2.86]
Baulac 2010	26	152	24	141	29.6%	1.00 [0.52 , 1.95]
Beydoun 2005	27	215	5	98	8.2%	2.46 [0.73 , 8.29]
Elger 2005	47	268	10	73	18.7%	1.28 [0.56 , 2.94]
French 2003	32	355	8	100	14.9%	1.13 [0.42 , 2.99]
French 2014	9	215	1	110	1.6%	4.60 [0.31 , 68.40]
Lee 2009	11	119	3	59	4.8%	1.82 [0.36 , 9.25]
Total (99% CI)		1716		772	100.0%	1.35 [0.94 , 1.93	1
Total events:	188		65				
Heterogeneity: Chi ² = 4	4.95, df = 7 (I	P = 0.67;	$[^2 = 0\%]$				0.01 0.1 1 10 100
Test for overall effect:	Z = 2.15 (P =	0.03)					Favours pregabalin Favours placebo
Trat for sub success differ							

Analysis 1.9. Comparison 1: Pregabalin versus placebo, Outcome 9: Fatigue

Test for subgroup differences: Not applicable

Analysis 1.10. Comparison 1: Pregabalin versus placebo, Outcome 10: Headache

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Antinew 2019	11	201	6	94	8.2%	0.86 [0.24 , 3.04]	
Arroyo 2004	17	191	15	97	20.0%	0.58 [0.25 , 1.35]	∣ _∎∔
Baulac 2010	18	152	28	141	29.2%	0.60 [0.29 , 1.22]	I _ ∎ ∔
Elger 2005	28	268	8	73	12.7%	0.95 [0.36 , 2.53]	I
French 2003	24	355	13	100	20.4%	0.52 [0.23 , 1.20]	Ⅰ _∎∔
Lee 2009	9	119	7	59	9.4%	0.64 [0.19 , 2.18]	· _•
Total (99% CI)		1286		564	100.0%	0.65 [0.45 , 0.94]	
Total events:	107		77				•
Heterogeneity: Chi ² = 2.	.04, df = 5 (F	P = 0.84); I	$I^2 = 0\%$				0.01 0.1 1 10 100
Test for overall effect: Z	L = 2.99 (P =	0.003)					Favours pregabalin Favours placebo
Test for subgroup differe	ences: Not aj	pplicable					

Analysis 1.11. Comparison 1: Pregabalin versus placebo, Outcome 11: Nausea

	Pregal	balin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Arroyo 2004	13	191	6	97	38.5%	1.10 [0.32 , 3.76]	
Beydoun 2005	11	215	6	98	39.9%	0.84 [0.23 , 2.97]	
Elger 2005	15	268	2	73	15.2%	2.04 [0.30 , 13.78]	
French 2014	4	215	1	110	6.4%	2.05 [0.12 , 35.88]	
Total (99% CI)		889		378	100.0%	1.20 [0.56 , 2.58]	
Total events:	43		15				
Heterogeneity: Chi ² = 1	1.32, df = 3 (F	P = 0.73);]	$I^2 = 0\%$			0.	101 0.1 1 10 100
Test for overall effect:	Z = 0.61 (P =	0.54)					ours pregabalin Favours placebo
Test for subgroup differ	rences: Not a	pplicable					

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	Pregal	oalin	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Antinew 2019	43	201	13	94	18.2%	1.55 [0.73 , 3.27]	_ _
Arroyo 2004	33	191	7	97	9.6%	2.39 [0.86 , 6.66]	
Baulac 2010	30	152	15	141	16.0%	1.86 [0.87 , 3.95]	
Beydoun 2005	57	215	12	98	17.0%	2.17 [1.02 , 4.61]	_ _
Elger 2005	49	268	6	73	9.7%	2.22 [0.77 , 6.43]	
French 2003	65	355	11	100	17.7%	1.66 [0.76 , 3.66]	+
French 2014	10	215	2	110	2.7%	2.56 [0.36 , 18.38]	
Lee 2009	26	119	3	59	4.1%	4.30 [0.94 , 19.57]	
Mann 2020	14	105	4	70	4.9%	2.33 [0.57 , 9.51]	
Total (99% CI)		1821		842	100.0%	2.05 [1.49 , 2.81]	•
Total events:	327		73				•
Heterogeneity: Chi ² = 3	8.46, df = 8 (H	P = 0.90); I	$1^2 = 0\%$			⊢ 0.0	1 0.1 1 10
Test for overall effect: 2	Z = 5.83 (P <	0.00001)				Favo	urs pregabalin Favours place
Test for subgroup differ	rences: Not a	pplicable					

Analysis 1.12. Comparison 1: Pregabalin versus placebo, Outcome 12: Somnolence

Analysis 1.13. Comparison 1: Pregabalin versus placebo, Outcome 13: Weight gain

	Pregal	oalin	Place	ebo		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI	
Antinew 2019	17	201	4	94	21.9%	1.99 [0.49 , 8.02]]	
Arroyo 2004	20	191	2	97	10.7%	5.08 [0.77 , 33.39]]	
Baulac 2010	14	152	2	141	8.3%	6.49 [0.95 , 44.46]]	
Beydoun 2005	38	215	2	98	11.0%	8.66 [1.37 , 54.65]]	
Elger 2005	53	268	5	73	31.6%	2.89 [0.91 , 9.17]]	
French 2003	20	355	0	100	3.1%	11.63 [0.29 , 459.09]]	
French 2014	8	215	0	110	2.7%	8.74 [0.21 , 366.41]]	
Lee 2009	14	119	2	59	10.7%	3.47 [0.52 , 23.28]]	_
Total (99% CI)		1716		772	100.0%	4.35 [2.34 , 8.11]		
Total events:	184		17				•	
Heterogeneity: Chi ² = 4	4.99, df = 7 (H	P = 0.66);]	$[^2 = 0\%]$				0.01 0.1 1 10	100
Test for overall effect:	Z = 6.09 (P <	0.00001)					Favours pregabalin Favours	s placebo
Test for subgroup differ	rences: Not a	pplicable						

Comparison 2. Pregabalin versus placebo - subgroup analysis

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.1 50% or greater reduction in seizure frequency - ITT	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.52, 2.12]
2.1.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 95% CI)	2.22 [1.36, 3.63]
2.1.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 95% CI)	2.86 [1.65, 4.94]

Pregabalin add-on for drug-resistant focal epilepsy (Review)



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.1.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 95% CI)	4.62 [3.34, 6.39]
2.1.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	1.76 [1.35, 2.30]
2.1.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.72, 1.48]
2.1.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	1.26 [0.91, 1.75]
2.2 50% or greater reduction in seizure frequency - best-case analysis	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.2.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 95% CI)	1.87 [1.03, 3.40]
2.2.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 95% CI)	3.18 [2.00, 5.06]
2.2.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 95% CI)	4.37 [2.61, 7.29]
2.2.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 95% CI)	7.72 [5.64, 10.57]
2.2.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	2.86 [2.24, 3.65]
2.2.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 95% CI)	1.28 [0.92, 1.79]
2.2.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	1.63 [1.21, 2.20]
2.3 50% or greater reduction in seizure frequency - worst-case analysis	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.3.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 95% CI)	0.55 [0.30, 0.99]
2.3.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.66, 1.38]
2.3.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 95% CI)	1.48 [0.98, 2.23]
2.3.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 95% CI)	1.72 [1.42, 2.09]
2.3.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.76, 1.12]
2.3.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 95% CI)	0.79 [0.57, 1.09]
2.3.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.72, 1.28]
2.4 Seizure freedom	4		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only

Pregabalin add-on for drug-resistant focal epilepsy (Review)



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
2.4.1 600 mg/d	2	523	Risk Ratio (M-H, Fixed, 95% CI)	6.92 [1.31, 36.70]	
2.4.2 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	2.39 [0.83, 6.89]	
2.5 Treatment withdrawal for any reason	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only	
2.5.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 95% CI)	0.87 [0.40, 1.89]	
2.5.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.41, 1.28]	
2.5.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 95% CI)	1.62 [0.85, 3.10]	
2.5.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 95% CI)	1.84 [1.42, 2.40]	
2.5.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	1.20 [0.89, 1.62]	
2.5.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 95% CI)	0.82 [0.36, 1.86]	
2.5.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [0.59, 2.46]	
2.6 Treatment withdrawal due to adverse effects	7		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only	
2.6.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 95% CI)	1.36 [0.43, 4.31]	
2.6.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.45, 2.32]	
2.6.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 95% CI)	2.89 [1.07, 7.78]	
2.6.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 95% CI)	3.78 [2.47, 5.81]	
2.6.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 95% CI)	2.26 [1.30, 3.95]	
2.6.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 95% CI)	1.09 [0.22, 5.27]	
2.6.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 95% CI)	2.57 [0.70, 9.45]	
2.7 Ataxia	6		Risk Ratio (M-H, Fixed, 99% Cl)	Subtotals only	
2.7.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	1.14 [0.14, 9.00]	
2.7.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	1.98 [0.56, 7.01]	
2.7.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	3.33 [0.62, 17.81]	
2.7.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 99% CI)	4.49 [2.25, 8.95]	

Pregabalin add-on for drug-resistant focal epilepsy (Review)



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.7.5 Titrated dose pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 99% CI)	4.46 [1.28, 15.48]
2.8 Dizziness	7		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.8.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	1.01 [0.31, 3.33]
2.8.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	2.04 [0.99, 4.22]
2.8.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	3.46 [1.39, 8.62]
2.8.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 99% CI)	3.72 [2.42, 5.69]
2.8.5 Titrated dose of pregabalin (150 to 600 mg/d pregabalin)	3	675	Risk Ratio (M-H, Fixed, 99% CI)	3.08 [1.80, 5.28]
2.8.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 99% CI)	5.99 [0.85, 42.02]
2.8.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 99% CI)	5.31 [0.76, 37.30]
2.9 Fatigue	7		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.9.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	0.71 [0.17, 2.94]
2.9.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	1.09 [0.50, 2.39]
2.9.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	1.53 [0.49, 4.76]
2.9.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 99% CI)	1.50 [0.89, 2.52]
2.9.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 99% CI)	1.15 [0.69, 1.91]
2.9.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 99% CI)	3.27 [0.17, 62.62]
2.9.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 99% CI)	5.79 [0.37, 91.55]
2.10 Headache	5		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.10.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	0.52 [0.16, 1.77]
2.10.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	0.53 [0.24, 1.17]
2.10.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	0.43 [0.12, 1.57]
2.10.4 600 mg/d pregabalin	3	588	Risk Ratio (M-H, Fixed, 99% CI)	0.63 [0.33, 1.19]
2.10.5 Titrated dose of pregabalin (150 to 600 mg/d)	3	675	Risk Ratio (M-H, Fixed, 99% CI)	0.74 [0.44, 1.25]

Pregabalin add-on for drug-resistant focal epilepsy (Review)



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.11 Nausea	4		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.11.1 150 mg/d pregabalin	1	196	Risk Ratio (M-H, Fixed, 99% CI)	1.31 [0.34, 5.00]
2.11.2 600 mg/d pregabalin	3	712	Risk Ratio (M-H, Fixed, 99% CI)	1.18 [0.51, 2.75]
2.11.3 Titrated dose of pregabalin (150 to 600 mg/d)	1	204	Risk Ratio (M-H, Fixed, 99% CI)	1.11 [0.12, 10.05]
2.11.4 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 99% CI)	3.27 [0.17, 62.62]
2.11.5 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 99% CI)	0.96 [0.03, 36.26]
2.12 Somnolence	7		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.12.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	0.93 [0.31, 2.78]
2.12.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	1.26 [0.58, 2.74]
2.12.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	1.62 [0.63, 4.12]
2.12.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 99% CI)	2.57 [1.64, 4.03]
2.12.5 Titrated dose of pregabalin	3	675	Risk Ratio (M-H, Fixed, 99% CI)	2.35 [1.31, 4.19]
2.12.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 99% CI)	2.18 [0.24, 19.70]
2.12.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 99% CI)	2.89 [0.36, 23.05]
2.13 Weight gain	7		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
2.13.1 50 mg/d pregabalin	1	188	Risk Ratio (M-H, Fixed, 99% CI)	3.40 [0.05, 224.69]
2.13.2 150 mg/d pregabalin	2	384	Risk Ratio (M-H, Fixed, 99% CI)	3.85 [0.64, 23.35]
2.13.3 300 mg/d pregabalin	1	190	Risk Ratio (M-H, Fixed, 99% CI)	14.43 [0.34, 620.87]
2.13.4 600 mg/d pregabalin	4	901	Risk Ratio (M-H, Fixed, 99% CI)	5.88 [2.52, 13.73]
2.13.5 Titrated dose of pregabalin (150 to 600 mg/d pregabalin)	3	675	Risk Ratio (M-H, Fixed, 99% CI)	3.64 [1.49, 8.87]
2.13.6 165 mg/d pregabalin (con- trolled release)	1	211	Risk Ratio (M-H, Fixed, 99% CI)	3.26 [0.05, 215.86]
2.13.7 330 mg/d pregabalin (con- trolled release)	1	224	Risk Ratio (M-H, Fixed, 99% CI)	14.48 [0.34, 613.54]

Pregabalin add-on for drug-resistant focal epilepsy (Review)

Analysis 2.1. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 1: 50% or greater reduction in seizure frequency - ITT

	Pregat	alin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
2.1.1 50 mg/d pregaba	lin						
French 2003		00	14	100	100.00/		
	13	88 88	14	100 100	100.0% 100.0%	1.06 [0.52 , 2.12]	
Subtotal (95% CI)	10	00	14	100	100.0%	1.06 [0.52 , 2.12]	
Total events:	13		14				
Heterogeneity: Not app Test for overall effect: 2		0.88)					
		,					
2.1.2 150 mg/d pregab			_				
Arroyo 2004	14	99	6	97	31.6%	2.29 [0.92 , 5.71]	
French 2003	27	88	14	100	68.4%	2.19 [1.23 , 3.91]	
Subtotal (95% CI)		187		197	100.0%	2.22 [1.36 , 3.63]	\bullet
Total events:	41		20				
Heterogeneity: Chi ² = 0			$^{2} = 0\%$				
Test for overall effect: 2	Z = 3.19 (P =	0.001)					
2.1.3 300 mg/d pregab	alin						
French 2003	36	90	14	100	100.0%	2.86 [1.65 , 4.94]	
Subtotal (95% CI)		90		100	100.0%	2.86 [1.65 , 4.94]	
Total events:	36		14				
Heterogeneity: Not app	licable						
Test for overall effect: 2		0.0002)					
2.1.4 600 mg/d pregab	alin						
Arroyo 2004	40	92	6	97	14.0%	7.03 [3.13 , 15.79]	
Beydoun 2005	98	215	9	98	29.6%	4.96 [2.62, 9.41]	
Elger 2005	62	137	8	73	25.0%	4.13 [2.09 , 8.15]	
French 2003	45	89	14	100	31.5%	3.61 [2.13 , 6.12]	
Subtotal (95% CI)		533		368	100.0%	4.62 [3.34 , 6.39]	
Total events:	245	000	37	500	10010 / 0		
Heterogeneity: Chi ² = 2		$= 0.57 \cdot 1$					
Test for overall effect: 2		-	070				
2.1.5 Titrated dose of J	aregahalin (1	50 to 600	mg/d)				
Baulac 2010	54	150 10 000	111g/U) 30	141	46.6%	1.67 [1.14 , 2.45]	
Elger 2005	41	131	8	73	15.4%	2.86 [1.42, 5.76]	■_
Lee 2009	55	119	19	59	38.0%	1.44 [0.94 , 2.18]	
Subtotal (95% CI)	55	402	15		100.0%	1.44 [0.94 , 2.10] 1.76 [1.35 , 2.30]	
Total events:	150	402	57	213	100.0 /0	1.70 [1.00 , 4.00]	
Heterogeneity: Chi ² = 2		$= 0.24) \cdot 1$					
Test for overall effect: 2			2370				
2.1.6 165 mg/d pregab	alin (contra-1	lad valaa-	a)				
2.1.6 165 mg/d pregab French 2014	alin (control 37		,	110	100 00/	1 03 [0 72 1 40]	
	3/	101	39	110	100.0%	1.03 [0.72 , 1.48]	—
Subtotal (95% CI)	25	101		110	100.0%	1.03 [0.72 , 1.48]	•
Total events:	37		39				
Heterogeneity: Not app Test for overall effect: 2		0.86)					
2 1 7 220 ma/d mas-L	alin (control	lad value -	c)				
2.1.7 330 mg/d pregab			·	110	100.007		
French 2014	51	114	39	110	100.0%	1.26 [0.91 , 1.75]	
Subtotal (95% CI)		114	39	110	100.0%	1.26 [0.91 , 1.75]	•
Total events:	51						

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.1. (Continued)

	-		 		7
Total events:	51	39			,
Heterogeneity: Not applicab	le				
Test for overall effect: $Z = 1$.41 (P = 0.16)				
				0.01 0.1 1	10 100
				Favours placebo	Favours pregabalin

Analysis 2.2. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 2: 50% or greater reduction in seizure frequency - best-case analysis

	Pregał	oalin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
2.2.1 50 mg/d pregaba	lin						
French 2003	23	88	14	100	100.0%	1.87 [1.03 , 3.40]	
Subtotal (95% CI)		88		100	100.0%	1.87 [1.03 , 3.40]	
Total events:	23		14	100	10010 / 0	107 [100 ; 01 10]	
Heterogeneity: Not app			11				
Test for overall effect: 2		0.04)					
2.2.2 150 mg/d pregab	alin						
Arroyo 2004	25	99	6	97	31.6%	4.08 [1.75, 9.51]	
French 2003	34	88	14	100	68.4%	2.76 [1.59 , 4.80]	
Subtotal (95% CI)		187		197	100.0%	3.18 [2.00 , 5.06]	
Total events:	59		20				
Heterogeneity: Chi ² = 0		P = 0.44): 1					
Test for overall effect: 2			0,0				
2.2.3 300 mg/d pregab	alin						
French 2003	55	90	14	100	100.0%	4.37 [2.61 , 7.29]	
Subtotal (95% CI)		90		100	100.0%	4.37 [2.61 , 7.29]	
Total events:	55		14	100		,	
Heterogeneity: Not app			- 1				
Test for overall effect: 2		0.00001)					
2.2.4 600 mg/d pregab	alin						
Arroyo 2004	63	92	6	97	14.0%	11.07 [5.04 , 24.33]	_
Beydoun 2005	157	215	9	98	29.6%	7.95 [4.24 , 14.90]	
Elger 2005	119	137	8	73	25.0%	7.93 [4.11 , 15.29]	
French 2003	73	89	14	100	31.5%	5.86 [3.57, 9.62]	
Subtotal (95% CI)		533		368	100.0%	7.72 [5.64 , 10.57]	
Total events:	412		37				
Heterogeneity: Chi ² = 2		P = 0.57: 1					
Test for overall effect: 2							
2.2.5 Titrated dose of	pregabalin (1	150 to 600) mg/d)				
Baulac 2010	100	152	30	141	46.6%	3.09 [2.21 , 4.33]	-
Elger 2005	72	131	8	73	15.4%	5.02 [2.56 , 9.82]	
Lee 2009	65	119	19	59	38.0%	1.70 [1.13 , 2.54]	
Subtotal (95% CI)		402		273	100.0%	2.86 [2.24 , 3.65]	
Total events:	237		57			-	•
Heterogeneity: Chi ² = 9 Test for overall effect: 7			I ² = 78%				
2.2.6 165 mg/d pregab			0)				
2.2.6 165 mg/d pregab French 2014	ann (control 46	101 releas	e) 39	110	100.0%		
	40	101 101	39		100.0% 100.0%	1.28 [0.92, 1.79]	
Subtotal (95% CI)	40	101	20	110	100.0%	1.28 [0.92 , 1.79]	•
Total events:	46		39				
Heterogeneity: Not app Test for overall effect: 2		0.14)					
2.2.7 330 mg/d pregab	alin (control	led releas	e)				
French 2014	66	114	ý 39	110	100.0%	1.63 [1.21 , 2.20]	
Subtotal (95% CI)		114		110	100.0%	1.63 [1.21 , 2.20]	
Total events:	66	-	39	-		. ,	
	licabla		00				

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.2. (Continued)

	-	- •	 	 -~,			
Total events:	66	39				•	
Heterogeneity: Not application	able						
Test for overall effect: Z =	3.24 (P = 0.001)						
				0.01	0.1	1 10	100
				Favor	ırs placebo	Favours	pregabalin

Analysis 2.3. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 3: 50% or greater reduction in seizure frequency - worst-case analysis

	Pregat	oalin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
2.3.1 50 mg/d pregaba	lin						
French 2003	13	88	27	100	100.0%	0.55 [0.30 , 0.99]	
Subtotal (95% CI)	15	88	27	100	100.0%	0.55 [0.30 , 0.99]	
Total events:	13	00	27	100	100.0 /0	0.55 [0.50 , 0.55]	
Heterogeneity: Not app			27				
Test for overall effect: 2		0.05)					
2.3.2 150 mg/d pregab	alin						
Arroyo 2004	14	99	19	97	43.2%	0.72 [0.38 , 1.36]	
French 2003	27	88	27	100	56.8%	1.14 [0.72 , 1.78]	
Subtotal (95% CI)	_/	187	_/	197	100.0%	0.96 [0.66 , 1.38]	
Total events:	41	107	46	157	100.0 /0	0.50 [0.00 ; 1.50]	—
Heterogeneity: Chi ² = 1		$0 = 0.25 \cdot 1$					
Test for overall effect:			1 - 2370				
2.3.3 300 mg/d pregab	alin						
French 2003	36	90	27	100	100.0%	1.48 [0.98 , 2.23]	
Subtotal (95% CI)	50	90 90	27	100 100	100.0% 100.0%	1.48 [0.98 , 2.23]	
Total events:	36	50	27	100	100.0 70	1.40 [0.30 , 2.23]	
Heterogeneity: Not app			27				
Test for overall effect: 2		0.06)					
rest for overall effect: A	с – 1.00 (Р =	0.00)					
2.3.4 600 mg/d pregab	alin						
Arroyo 2004	40	92	19	97	16.5%	2.22 [1.39 , 3.54]	_
Beydoun 2005	98	215	26	98	31.8%	1.72 [1.20 , 2.46]	_
Elger 2005	62	137	25	73	29.1%	1.32 [0.92 , 1.91]	+ - -
French 2003	45	89	27	100	22.7%	1.87 [1.28 , 2.74]	_
Subtotal (95% CI)		533		368	100.0%	1.72 [1.42 , 2.09]	
Total events:	245		97				•
Heterogeneity: Chi ² = 3	3.32, df = 3 (P	e = 0.34); I	[2 = 10%				
Test for overall effect: 2	Z = 5.48 (P <	0.00001)					
2.3.5 Titrated dose of	pregabalin (1	L50 to 600) mg/d)				
Baulac 2010	54	152	65	141	52.8%	0.77 [0.58 , 1.02]	
Elger 2005	41	131	25	73	25.2%	0.91 [0.61 , 1.37]	_ _
Lee 2009	55	119	21	59	22.0%	1.30 [0.88 , 1.93]	+- -
Subtotal (95% CI)		402		273	100.0%	0.92 [0.76 , 1.12]	
Total events:	150		111				٦
Heterogeneity: Chi ² = 4 Test for overall effect: 2			2 = 55%				
		ŗ					
2.3.6 165 mg/d pregab French 2014	alin (control 37	led releas 101	e) 51	110	100.0%	0.79 [0.57 , 1.09]	
Subtotal (95% CI)	57	101	51	110	100.0%	0.79 [0.57 , 1.09]	
Total events:	37	101	51	110	100.0 /0	0.70 [0.07 , 1.00]	
Heterogeneity: Not app			51				
Test for overall effect: 2		0.16)					
2.3.7 330 mg/d pregab	alin (control	led releas	e)				
French 2014	51	114	51	110	100.0%	0.96 [0.72 , 1.28]	
Subtotal (95% CI)	51	114	51	110	100.0%	0.96 [0.72 , 1.28]	
Total events:	51	114	51	110	100.0 /0	0.00 [0.72 , 1.20]	\blacksquare
rour crento.	51		51				

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.3. (Continued)



Analysis 2.4. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 4: Seizure freedom

	Pregal	Pregabalin		Placebo		Risk Ratio	Risk Ratio		
Study or Subgroup	Events	Total	Total Events		Weight	M-H, Fixed, 95% CI	M-H, Fixed	, 95% CI	
2.4.1 600 mg/d									
Beydoun 2005	17	215	0	98	34.4%	16.04 [0.97 , 264.07]			
Elger 2005	4	137	1	73	65.6%	2.13 [0.24 , 18.72]			
Subtotal (95% CI)		352		171	100.0%	6.92 [1.31 , 36.70]			
Total events:	21		1						
Heterogeneity: Chi ² = 1	.48, df = 1 (I	P = 0.22);]	I² = 32%						
Test for overall effect: 2	Z = 2.27 (P =	0.02)							
2.4.2 Titrated dose of J	pregabalin (150 to 600) mg/d)						
Baulac 2010	6	152	1	141	20.8%	5.57 [0.68 , 45.66]	+	_	
Elger 2005	4	131	1	73	25.7%	2.23 [0.25 , 19.57]			
Lee 2009	5	119	2	59	53.5%	1.24 [0.25 , 6.20]		<u> </u>	
Subtotal (95% CI)		402		273	100.0%	2.39 [0.83 , 6.89]			
Total events:	15		4					•	
Heterogeneity: Chi ² = 1	.26, df = 2 (I	P = 0.53); I	$I^2 = 0\%$						
Test for overall effect: 2	Z = 1.62 (P =	0.11)							
							0.005 0.1 1 Favours placebo	10 200 Favours pregabal	

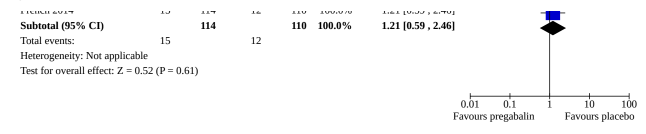
	Pregabalin		Placebo		Risk Ratio		Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
2.5.1 50 mg/d pregaba	lin							
French 2003	10	88	13	100	100.0%	0.87 [0.40 , 1.89]		
Subtotal (95% CI)		88		100	100.0%	0.87 [0.40 , 1.89]		
Total events:	10		13					
Heterogeneity: Not app	licable							
Test for overall effect: 2	Z = 0.34 (P =	0.73)						
2.5.2 150 mg/d pregab	alin							
Arroyo 2004	11	99	13	97	51.9%	0.83 [0.39 , 1.76]		
French 2003	7	88	13	100	48.1%			
Subtotal (95% CI)		187	-	197				
Total events:	18	107	26	107	10000 /0	0		
Heterogeneity: $Chi^2 = 0$		P = 0.61): 1						
Test for overall effect: 2			0,0					
2.5.3 300 mg/d pregab	alin							
French 2003	19	90	13	100	100.0%	1.62 [0.85 , 3.10]		
Subtotal (95% CI)	15	90	15	100	100.0%			
Total events:	19	50	13	100	100.070	1.02 [0.00 ; 0.10]		
Heterogeneity: Not app			15					
Test for overall effect: 2		0.14)						
7 5 4 6 0 0 1 1 1 1 1 1 1 1 1 1	-14							
2.5.4 600 mg/d pregab		02	10	07	10.00/	107[101 346]		
Arroyo 2004	23	92	13	97	18.0%			
Beydoun 2005	59	215	17	98 72	33.2%			
Elger 2005	57	137	17	73	31.5%			
French 2003	28	89	13	100	17.4%			
Subtotal (95% CI)	407	533	60	368	100.0%	1.84 [1.42 , 2.40]		
Total events:	167		60					
Heterogeneity: $Chi^2 = 1$			$l^2 = 0\%$					
Test for overall effect: 2	Z = 4.56 (P <	0.00001)						
2.5.5 Titrated dose of J			•					
Baulac 2010	46	152	35	141	59.7%		₽	
Elger 2005	31	131	17	73	35.9%		-+-	
Lee 2009	10	119	2	59				
Subtotal (95% CI)	^ =	402		273	100.0%	1.20 [0.89 , 1.62]	•	
Total events:	87		54					
Heterogeneity: Chi ² = 1 Test for overall effect: 2			L ² = 0%					
2 E 6 16E mg/d mas-L	alin (contral	lad valar -	a)					
2.5.6 165 mg/d pregab French 2014	alın (control 9	led releas 101	e) 12	110	100.0%	0.82 [0.36 , 1.86]		
Subtotal (95% CI)		101		110	100.0%	0.82 [0.36 , 1.86]		
Total events:	9		12			-	\mathbf{T}	
Heterogeneity: Not app								
Test for overall effect: 2		0.63)						
2.5.7 330 mg/d pregab	alin (control	led releas	e)					
French 2014	15	114	12	110	100.0%	1.21 [0.59 , 2.46]	_	
Subtotal (95% CI)	-	114		110			_	
- '	· -	•		0			$\mathbf{\mathbf{T}}$	

Analysis 2.5. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 5: Treatment withdrawal for any reason

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.5. (Continued)



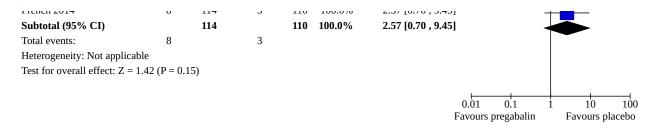
Analysis 2.6. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 6: Treatment withdrawal due to adverse effects

	Pregat	alin	Placebo		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
2.6.1 50 mg/d pregabal	lin						
French 2003	6	88	5	100	100.0%	1.36 [0.43 , 4.31]	
Subtotal (95% CI)	0	88	5	100	100.0%	1.36 [0.43 , 4.31]	
Total events:	6	00	5	100	100.070	1.50 [0.45 ; 4.51]	
Heterogeneity: Not appl			5				
Test for overall effect: Z		0.60)					
2.6.2 150 mg/d pregaba	alin						
Arroyo 2004	10	99	6	97	56.4%	1.63 [0.62 , 4.32]	_ _
French 2003	1	88	5	100	43.6%	0.23 [0.03 , 1.91]	
Subtotal (95% CI)		187		197	100.0%	1.02 [0.45 , 2.32]	-
Total events:	11		11				
Heterogeneity: Chi ² = 2.	.81, df = 1 (P	= 0.09); 1	$^{2} = 64\%$				
Test for overall effect: Z							
2.6.3 300 mg/d pregaba	alin						
French 2003	13	90	5	100	100.0%	2.89 [1.07 , 7.78]	
Subtotal (95% CI)		90		100	100.0%	2.89 [1.07 , 7.78]	
Total events:	13		5				
Heterogeneity: Not appl	licable						
Test for overall effect: Z		0.04)					
2.6.4 600 mg/d pregaba	alin						
Arroyo 2004	17	92	6	97	21.9%	2.99 [1.23 , 7.24]	
Beydoun 2005	48	215	7	98	36.0%	3.13 [1.47 , 6.66]	_
Elger 2005	45	137	5	73	24.4%	4.80 [1.99 , 11.55]	
French 2003	21	89	5	100	17.6%	4.72 [1.86 , 11.99]	
Subtotal (95% CI)		533		368	100.0%	3.78 [2.47 , 5.81]	
_							
Total events:	131		23				
Heterogeneity: Chi ² = 1.	.01, df = 3 (P						
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p	.01, df = 3 (P Z = 6.09 (P < Dregabalin (1	0.00001) 1 50 to 600	² = 0% mg/d)	4.44	E0.404	2 22 [1 40 4 40]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24	0.00001) 1 50 to 600 152	² = 0% mg/d) 10	141	59.4%	2.23 [1.10 , 4.49]	-
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16	0.00001) 1 50 to 600 152 131	² = 0% mg/d) 10 5	73	36.8%	1.78 [0.68 , 4.67]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24	0.00001) 50 to 600 152 131 119	² = 0% mg/d) 10	73 59	36.8% 3.8%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI)	.01, df = 3 (P Z = 6.09 (P < oregabalin (1 24 16 7	0.00001) 1 50 to 600 152 131	² = 0% mg/d) 10 5 0	73 59	36.8%	1.78 [0.68 , 4.67]	
Total events: Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events:	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 47	0.00001) 50 to 600 152 131 119 402	² = 0% mg/d) 10 5 0 15	73 59	36.8% 3.8%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI)	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 47 .92, df = 2 (P	0.00001) 150 to 600 152 131 119 402 9 = 0.63); 1	² = 0% mg/d) 10 5 0 15	73 59	36.8% 3.8%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13]	
Heterogeneity: $Chi^2 = 1$. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: $Chi^2 = 0$. Test for overall effect: Z	.01, df = 3 (P Z = 6.09 (P < bregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P =	0.00001) 150 to 600 152 131 119 402 1 = 0.63); 1 0.004)	$2^{2} = 0\%$ mg/d) 10 5 0 15 $2^{2} = 0\%$	73 59	36.8% 3.8%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P = alin (control	0.00001) 150 to 600 152 131 119 402 -= 0.63); 1 0.004) led releas	mg/d) mg/d) 10 5 0 15 $^{2} = 0\%$ e)	73 59 273	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014	.01, df = 3 (P Z = 6.09 (P < bregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P =	0.00001) 150 to 600 152 131 119 402 = 0.63); 1 0.004) led releas 101	$2^{2} = 0\%$ mg/d) 10 5 0 15 $2^{2} = 0\%$	73 59 273 110	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95] 1.09 [0.22 , 5.27]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014 Subtotal (95% CI)	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P = alin (control 3	0.00001) 150 to 600 152 131 119 402 -= 0.63); 1 0.004) led releas	mg/d) mg/d) 10 5 0 15 $^{2} = 0\%$ 15 a a a b 3	73 59 273	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014 Subtotal (95% CI) Total events:	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P = alin (control 3 3	0.00001) 150 to 600 152 131 119 402 = 0.63); 1 0.004) led releas 101	mg/d) mg/d) 10 5 0 15 $^{2} = 0\%$ e)	73 59 273 110	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95] 1.09 [0.22 , 5.27]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014 Subtotal (95% CI) Total events: Heterogeneity: Not appl	0.01, df = 3 (P Z = 6.09 (P < pregabalin (1) 24 16 7 47 .92, df = 2 (P Z = 2.89 (P = alin (control) 3 3 licable	0.00001) 150 to 600 152 131 119 402 1 = 0.63); 1 0.004) led releas 101 101	mg/d) mg/d) 10 5 0 15 $^{2} = 0\%$ 15 a a a b 3	73 59 273 110	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95] 1.09 [0.22 , 5.27]	
Heterogeneity: $Chi^2 = 1$. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: $Chi^2 = 0$. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014 Subtotal (95% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P = alin (control 3 3 Licable Z = 0.11 (P =	0.00001) 150 to 600 152 131 119 402 1 = 0.63); 1 0.004) 101 101 0.92)	$2^{2} = 0\%$ mg/d) 10 5 0 15 $2^{2} = 0\%$ r r r r r r r r	73 59 273 110	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95] 1.09 [0.22 , 5.27]	
Heterogeneity: Chi ² = 1. Test for overall effect: Z 2.6.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (95% CI) Total events: Heterogeneity: Chi ² = 0. Test for overall effect: Z 2.6.6 165 mg/d pregaba French 2014 Subtotal (95% CI) Total events: Heterogeneity: Not appl	.01, df = 3 (P Z = 6.09 (P < pregabalin (1 24 16 7 .92, df = 2 (P Z = 2.89 (P = alin (control 3 3 Licable Z = 0.11 (P =	0.00001) 150 to 600 152 131 119 402 1 = 0.63); 1 0.004) 101 101 0.92)	$2^{2} = 0\%$ mg/d) 10 5 0 15 $2^{2} = 0\%$ r r r r r r r r	73 59 273 110	36.8% 3.8% 100.0%	1.78 [0.68 , 4.67] 7.50 [0.44 , 129.13] 2.26 [1.30 , 3.95] 1.09 [0.22 , 5.27]	

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.6. (Continued)



	Pregal	Pregabalin Events Total		Placebo Events Total		Risk Ratio	Risk Ratio M-H, Fixed, 99% CI	
Study or Subgroup	Events					M-H, Fixed, 99% CI		
2.7.1 50 mg/d pregaba	alin							
French 2003	3	88	3	100	100.0%	1.14 [0.14 , 9.00]		
Subtotal (99% CI)		88		100	100.0%	1.14 [0.14 , 9.00]		
Total events:	3		3					
Heterogeneity: Not app	olicable							
Test for overall effect:	Z = 0.16 (P =	0.87)						
2.7.2 150 mg/d pregab	oalin							
Arroyo 2004	2	99	3	97	51.9%	0.65 [0.06 , 6.66]		
French 2003	9	88	3	100	48.1%	3.41 [0.64 , 18.21]	-	
Subtotal (99% CI)		187		197	100.0%	1.98 [0.56 , 7.01]		
Fotal events:	11		6					
Heterogeneity: Chi ² = 2		P = 0.14):						
Test for overall effect:								
2.7.3 300 mg/d pregab	oalin							
French 2003	9	90	3	100	100.0%	3.33 [0.62 , 17.81]		
Subtotal (99% CI)		90	-	100	100.0%	3.33 [0.62 , 17.81]		
Fotal events:	9		3			,		
Heterogeneity: Not app								
Test for overall effect:		0.06)						
2.7.4 600 mg/d pregab	oalin							
Arroyo 2004	16	92	3	97	16.3%	5.62 [1.16, 27.21]		
Beydoun 2005	48	215	6	98	46.0%	3.65 [1.25 , 10.63]		
Elger 2005	29	137	3	73	21.9%	5.15 [1.13 , 23.48]		
French 2003	13	89	3	100	15.8%	4.87 [0.98 , 24.28]		
Subtotal (99% CI)		533		368	100.0%	4.49 [2.25 , 8.95]		
Total events:	106		15					
Heterogeneity: Chi ² = (P = 0.93);]						
Test for overall effect:		· · ·						
2.7.5 Titrated dose pro	egabalin (150) to 600 m	ıg/d)					
- Jaulac 2010	13	152	1	141	18.7%	12.06 [0.85 , 171.72]	_	
Elger 2005	12	131	3	73	69.3%	2.23 [0.44 , 11.26]		
Lee 2009	5	119	0	59	12.0%	5.50 [0.13 , 241.69]		
Subtotal (99% CI)		402		273	100.0%	4.46 [1.28 , 15.48]		
Fotal events:	30		4			- / 1		
Heterogeneity: Chi ² = 2		P = 0.34): 1						
Test for overall effect:								
		····-)						
						⊢ 0.0	1 0.1 1 10 1	
							urs pregabalin Favours placel	

Analysis 2.7. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 7: Ataxia

Analysis 2.8. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 8: Dizziness

	Pregab	alin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% C
2.8.1 50 mg/d pregabal	lin						
French 2003	8	88	9	100	100.0%	1.01 [0.31 , 3.33]	
Subtotal (99% CI)		88		100	100.0%	1.01 [0.31 , 3.33]	
Total events:	8		9				
Heterogeneity: Not appl							
Test for overall effect: Z		0.98)					
2.8.2 150 mg/d pregaba	alin						
Arroyo 2004	19	99	8	97	49.0%	2.33 [0.84 , 6.46]	
French 2003	14	88	9	100	51.0%	1.77 [0.63 , 4.97]	
Subtotal (99% CI)		187		197	100.0%	2.04 [0.99 , 4.22]	
Total events:	33		17			- / -	
Heterogeneity: $Chi^2 = 0$		= 0.63); I	$^{2} = 0\%$				
Test for overall effect: Z							
2.8.3 300 mg/d pregaba	alin						
French 2003	28	90	9	100	100.0%	3.46 [1.39 , 8.62]	
Subtotal (99% CI)		90		100	100.0%	3.46 [1.39 , 8.62]	
Total events:	28		9				
Heterogeneity: Not appl	licable						
Test for overall effect: Z	Z = 3.50 (P = 0)	0.0005)					
2.8.4 600 mg/d pregaba	alin						
Arroyo 2004	24	92	8	97	18.0%	3.16 [1.18 , 8.45]	
Beydoun 2005	88	215	14	98	44.4%	2.87 [1.46 , 5.61]	
Elger 2005	59	137	6	73	18.1%	5.24 [1.85 , 14.81]	
French 2003	38	89	9	100	19.6%	4.74 [1.97 , 11.41]	
Subtotal (99% CI)		533		368	100.0%	3.72 [2.42 , 5.69]	•
Total events:	209		37				•
Heterogeneity: Chi ² = 2		= 0.49); I	$^{2} = 0\%$				
Test for overall effect: Z	Z = 7.92 (P <	0.00001)					
2.8.5 Titrated dose of p		150 to 600	mg/d preg	abalin)			
2.8.5 Titrated dose of p Baulac 2010	pregabalin (1 38	1 50 to 600 152	13	141	46.2%	2.71 [1.25 , 5.86]	
2.8.5 Titrated dose of p Baulac 2010 Elger 2005	pregabalin (1 38 32	1 50 to 600 152 131	13 6	141 73	26.4%	2.97 [1.01 , 8.77]	
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009	pregabalin (1 38	1 50 to 600 152 131 119	13	141 73 59	26.4% 27.5%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76]	
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI)	pregabalin (1 38 32 46	1 50 to 600 152 131	13 6 6	141 73 59	26.4%	2.97 [1.01 , 8.77]	• •
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events:	pregabalin (1 38 32 46 116	150 to 600 152 131 119 402	13 6 6 25	141 73 59	26.4% 27.5%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76]	•
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI)	pregabalin (1 38 32 46 116 0.46, df = 2 (P	150 to 600 152 131 119 402 = 0.79); I	13 6 6 25	141 73 59	26.4% 27.5%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76]	•
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0	150 to 600 152 131 119 402 = 0.79); I 0.00001)	13 6 6 25 $^{2} = 0\%$	141 73 59	26.4% 27.5%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76]	•
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0	150 to 600 152 131 119 402 = 0.79); I 0.00001)	13 6 6 25 $^{2} = 0\%$	141 73 59	26.4% 27.5%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76]	•
 2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi² = 0 Test for overall effect: 2 2.8.6 165 mg/d pregaba French 2014 	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll	150 to 600 152 131 119 402 = 0.79); I 0.00001) led release	13 6 6 25 ² = 0%	141 73 59 273	26.4% 27.5% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28]	•
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.8.6 165 mg/d pregaba	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll	150 to 600 152 131 119 402 = 0.79); I 0.00001) led releas 101	13 6 6 25 ² = 0%	141 73 59 273 110	26.4% 27.5% 100.0% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28] 5.99 [0.85 , 42.02]	•
 2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi² = 0 Test for overall effect: Z 2.8.6 165 mg/d pregaba French 2014 Subtotal (99% CI) Total events: 	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll 11	150 to 600 152 131 119 402 = 0.79); I 0.00001) led releas 101	13 6 25 $2^2 = 0\%$ e) 2	141 73 59 273 110	26.4% 27.5% 100.0% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28] 5.99 [0.85 , 42.02]	•
2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: 2 2.8.6 165 mg/d pregaba French 2014 Subtotal (99% CI)	pregabalin (1 38 32 46 116 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll 11 11 11	150 to 600 152 131 119 402 = 0.79); I 0.00001) led releas 101 101	13 6 25 $2^2 = 0\%$ e) 2	141 73 59 273 110	26.4% 27.5% 100.0% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28] 5.99 [0.85 , 42.02]	
 2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi² = 0 Test for overall effect: Z 2.8.6 165 mg/d pregaba French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl 	pregabalin (1 38 32 46 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll 11 11 11 Licable Z = 2.37 (P = 0	150 to 600 152 131 119 402 = 0.79); I 0.00001) led releas 101 101 0.02)	$ \begin{array}{c} 13\\ 6\\ 25\\ 2=0\%\\ e) 2\\ 2\\ 2 \end{array} $	141 73 59 273 110	26.4% 27.5% 100.0% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28] 5.99 [0.85 , 42.02]	
 2.8.5 Titrated dose of p Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi² = 0 Test for overall effect: Z 2.8.6 165 mg/d pregaba French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z 	pregabalin (1 38 32 46 0.46, df = 2 (P Z = 5.37 (P < 0 alin (controll 11 11 11 Licable Z = 2.37 (P = 0	150 to 600 152 131 119 402 = 0.79); I 0.00001) led releas 101 101 0.02)	$ \begin{array}{c} 13\\ 6\\ 25\\ 2=0\%\\ e) 2\\ 2\\ 2 \end{array} $	141 73 59 273 110	26.4% 27.5% 100.0% 100.0%	2.97 [1.01 , 8.77] 3.80 [1.34 , 10.76] 3.08 [1.80 , 5.28] 5.99 [0.85 , 42.02]	

Pregabalin add-on for drug-resistant focal epilepsy (Review)

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Analysis 2.8. (Continued)

11011011 2014	11	114	4	110	100.070	J.JT [0./0 , J/.J(ני		+		-
Subtotal (99% CI)		114		110	100.0%	5.31 [0.76 , 37.30]				-
Total events:	11		2								
Heterogeneity: Not applica	able										
Test for overall effect: Z =	2.20 (P = 0.0)3)									
							0.01	0.1	1	10	100
							Favours pre	egabalin	Favo	ours pl	acebo

Analysis 2.9. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 9: Fatigue

	Pregat	Pregabalin		Placebo		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI	
2.9.1 50 mg/d pregaba	lin							
French 2003	5	88	8	100	100.0%	0.71 [0.17 , 2.94]		
Subtotal (99% CI)		88		100	100.0%	0.71 [0.17 , 2.94]		
Total events:	5		8					
Heterogeneity: Not app			-					
Test for overall effect: 2		0.53)						
2.9.2 150 mg/d pregab	alin							
Arroyo 2004	13	99	11	97	59.7%	1.16 [0.43 , 3.11]		
French 2003	7	88	8	100	40.3%	0.99 [0.28 , 3.57]		
Subtotal (99% CI)		187		197	100.0%	1.09 [0.50 , 2.39]		
Total events:	20		19			. , .		
Heterogeneity: $Chi^2 = 0$).06, df = 1 (F	P = 0.81;	$2^{2} = 0\%$					
Test for overall effect: 2		· · ·						
2.9.3 300 mg/d pregab	alin							
French 2003	11	90	8	100	100.0%	1.53 [0.49 , 4.76]		
Subtotal (99% CI)		90		100	100.0%	1.53 [0.49 , 4.76]		
Total events:	11		8					
Heterogeneity: Not app	licable							
Test for overall effect: 2	Z = 0.96 (P =	0.34)						
2.9.4 600 mg/d pregab	alin							
Arroyo 2004	13	92	11	97	28.1%	1.25 [0.46 , 3.34]	_	
Beydoun 2005	27	215	5	98	18.0%	2.46 [0.73 , 8.29]		
Elger 2005	25	137	10	73	34.2%	1.33 [0.55 , 3.24]	_	
French 2003	9	89	8	100	19.7%	1.26 [0.38 , 4.17]		
Subtotal (99% CI)		533		368	100.0%	1.50 [0.89 , 2.52]	•	
Total events:	74		34				•	
Heterogeneity: Chi ² = 1 Test for overall effect: 2			$1^2 = 0\%$					
2.9.5 Titrated dose of j	pregabalin (1	150 to 600	mg/d)					
Baulac 2010	26	152	24	141	59.6%	1.00 [0.52 , 1.95]		
Elger 2005	22	131	10	73	30.8%	1.23 [0.49 , 3.04]	_ _	
Lee 2009	11	119	3	59	9.6%	1.82 [0.36 , 9.25]	_	
Subtotal (99% CI)		402		273	100.0%	1.15 [0.69 , 1.91]	•	
Total events:	59		37					
Heterogeneity: Chi ² = 0 Test for overall effect: 2		,	$1^2 = 0\%$					
2.9.6 165 mg/d pregab				110	100.00/			
French 2014	3	101	1	110	100.0%	3.27 [0.17, 62.62]		
Subtotal (99% CI)	~	101		110	100.0%	3.27 [0.17 , 62.62]		
Total events:	3		1					
Heterogeneity: Not app	licable Z = 1.03 (P =	0.30)						
rest for overall effect. 2								
	alin (control	led releas	e)					
2.9.7 330 mg/d pregab French 2014	alin (control 6	led releas 114	e) 1	110	100.0%	5.79 [0.37 , 91.55]		

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.9. (Continued)

1101011 201 4	U	114	T	110	100.0 /0	J./J [U.J/ , JI.JJ		
Subtotal (99% CI)		114		110	100.0%	5.79 [0.37 , 91.55		
Total events:	6		1					
Heterogeneity: Not applica	able							
Test for overall effect: Z =	1.64 (P = 0.10	0)						
							0.01 0.1	1 10 100
							Favours pregabalin	Favours placebo

Pregabalin Placebo **Risk Ratio Risk Ratio** M-H, Fixed, 99% CI Study or Subgroup Events Total Events Total Weight M-H, Fixed, 99% CI 2.10.1 50 mg/d pregabalin French 2003 88 13 100.0% 0.52 [0.16, 1.77] 6 100 Subtotal (99% CI) 88 100.0% 0.52 [0.16 , 1.77] 100 Total events: 6 13 Heterogeneity: Not applicable Test for overall effect: Z = 1.37 (P = 0.17)2.10.2 150 mg/d pregabalin 0.39 [0.12, 1.29] Arroyo 2004 6 99 15 97 55.5% 88 French 2003 8 13 100 44.5% 0.70 [0.23 , 2.09] Subtotal (99% CI) 187 197 100.0% 0.53 [0.24 , 1.17] Total events: 14 28 Heterogeneity: Chi² = 0.85, df = 1 (P = 0.36); I² = 0% Test for overall effect: Z = 2.06 (P = 0.04)2.10.3 300 mg/d pregabalin French 2003 5 90 13 100 100.0% 0.43 [0.12 , 1.57] Subtotal (99% CI) 90 100 100.0% 0.43 [0.12 , 1.57] 5 13 Total events: Heterogeneity: Not applicable Test for overall effect: Z = 1.68 (P = 0.09)2.10.4 600 mg/d pregabalin 92 15 97 39.2% 0.77 [0.30, 2.00] Arroyo 2004 11 0.67 [0.21 , 2.13] Elger 2005 10 137 8 28.0% 73 French 2003 5 89 32.8% 0.43 [0.12 , 1.59] 13 100 Subtotal (99% CI) 100.0% 318 270 0.63 [0.33 , 1.19] 26 Total events: 36 Heterogeneity: $Chi^2 = 0.88$, df = 2 (P = 0.64); $I^2 = 0\%$ Test for overall effect: Z = 1.86 (P = 0.06)2.10.5 Titrated dose of pregabalin (150 to 600 mg/d) Baulac 2010 18 152 28 141 59.7% 0.60 [0.29 , 1.22] Elger 2005 18 131 8 73 21.1% 1.25 [0.45 , 3.50] Lee 2009 9 119 7 59 19.2% 0.64 [0.19 , 2.18] Subtotal (99% CI) 402 273 100.0% 0.74 [0.44 , 1.25] Total events: 45 43 Heterogeneity: Chi² = 2.45, df = 2 (P = 0.29); I² = 18% Test for overall effect: Z = 1.46 (P = 0.14)

0.01

0.1

Favours pregabalin

Analysis 2.10. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 10: Headache

100

10

Favours placebo

Analysis 2.11. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 11: Nausea

	Pregab	alin	Place	ebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
2.11.1 150 mg/d pregaba	lin						
Arroyo 2004	8	99	6	97	100.0%	1.31 [0.34 , 5.00]	
Subtotal (99% CI)		99		97	100.0%	1.31 [0.34 , 5.00]	
Total events:	8		6				
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 0.51 (P =	0.61)					
2.11.2 600 mg/d pregaba	lin						
Arroyo 2004	5	92	6	97	35.0%	0.88 [0.19 , 3.99]	
3eydoun 2005	11	215	6	98	49.4%	0.84 [0.23 , 2.97]	
Elger 2005	11	137	2	73	15.6%	2.93 [0.42 , 20.49]	
Subtotal (99% CI)		444		268	100.0%	1.18 [0.51 , 2.75]	
Total events:	27		14				\mathbf{T}
Heterogeneity: Chi ² = 2.1	9, df = 2 (P	= 0.33); I	$1^2 = 9\%$				
Test for overall effect: Z =	= 0.50 (P =	0.62)					
2.11.3 Titrated dose of p	regabalin (150 to 60	0 mg/d)				
Elger 2005	4	131	2	73	100.0%	1.11 [0.12 , 10.05]	
Subtotal (99% CI)		131		73	100.0%	1.11 [0.12 , 10.05]	
Total events:	4		2				
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 0.13 (P =	0.90)					
2.11.4 165 mg/d pregaba	lin (contro	lled relea	se)				
French 2014	3	101	. 1	110	100.0%	3.27 [0.17 , 62.62]	
Subtotal (99% CI)		101		110	100.0%	3.27 [0.17 , 62.62]	
Total events:	3		1				
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 1.03 (P =	0.30)					
2.11.5 330 mg/d pregaba	lin (contro	lled relea	se)				
French 2014	1	114	1	110	100.0%	0.96 [0.03 , 36.26]	
Subtotal (99% CI)		114		110	100.0%	0.96 [0.03 , 36.26]	
Total events:	1		1				
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 0.03 (P =	0.98)					
						0.0	
						Favo	ours pregabalin Favours place

Analysis 2.12. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 12: Somnolence

	Pregat	oalin	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
2.12.1 50 mg/d pregab	alin						
French 2003	9	88	11	100	100.0%	0.93 [0.31 , 2.78]	
Subtotal (99% CI)	5	88		100	100.0%	0.93 [0.31 , 2.78]	
Total events:	9		11				
Heterogeneity: Not app							
Test for overall effect:		0.86)					
2.12.2 150 mg/d prega	balin						
Arroyo 2004	6	99	7	97	40.7%	0.84 [0.21 , 3.35]	
French 2003	15	88	11	100	59.3%	1.55 [0.60 , 4.01]	
Subtotal (99% CI)		187		197	100.0%	1.26 [0.58 , 2.74]	—
Total events:	21		18				
Heterogeneity: Chi ² = ().88, df = 1 (P	e = 0.35); 1	$2^{2} = 0\%$				
Test for overall effect:							
2.12.3 300 mg/d prega	balin						
French 2003	16	90	11	100	100.0%	1.62 [0.63 , 4.12]	_
Subtotal (99% CI)		90		100	100.0%	1.62 [0.63 , 4.12]	
Total events:	16		11				
Heterogeneity: Not app	licable						
Test for overall effect:	Z = 1.32 (P =	0.19)					
2.12.4 600 mg/d prega	balin						
Arroyo 2004	27	92	7	97	16.4%	4.07 [1.46 , 11.35]	
Beydoun 2005	57	215	12	98	39.7%	2.17 [1.02 , 4.61]	_ _
Elger 2005	24	137	6	73	18.9%	2.13 [0.70 , 6.50]	+
French 2003	25	89	11	100	25.0%	2.55 [1.09 , 5.99]	
Subtotal (99% CI)		533		368	100.0%	2.57 [1.64 , 4.03]	
Total events:	133		36				•
Heterogeneity: Chi ² = 1 Test for overall effect: 1			$2^2 = 0\%$				
2.12.5 Titrated dose of		150	15	1 4 1	F7.00/		
Baulac 2010	30 25	152	15	141	57.0%	1.86 [0.87 , 3.95]	+
				70	20 20/		
Elger 2005		131	6	73 E0	28.2%	2.32 [0.77 , 7.04]	+
Lee 2009	25 26	119	6 3	59	14.7%	4.30 [0.94 , 19.57]	
Lee 2009 Subtotal (99% CI)	26		3	59			•
Lee 2009 Subtotal (99% CI) Total events:	26 81	119 402	3 24	59	14.7%	4.30 [0.94 , 19.57]	•
Lee 2009 Subtotal (99% CI)	26 81 1.70, df = 2 (P	119 402 9 = 0.43); 1	3 24	59	14.7%	4.30 [0.94 , 19.57]	•
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1	26 81 1.70, df = 2 (P Z = 3.79 (P =	119 402 9 = 0.43);] 0.0001)	3 24 2 ² = 0%	59	14.7%	4.30 [0.94 , 19.57]	•
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1	26 81 1.70, df = 2 (P Z = 3.79 (P =	119 402 9 = 0.43);] 0.0001)	3 24 2 ² = 0%	59	14.7%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19]	•
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega French 2014	26 81 1.70, df = 2 (P Z = 3.79 (P = balin (contro	119 402 9 = 0.43); 1 0.0001) blled relea 101	3 24 2 ² = 0%	59 273 110	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19] 2.18 [0.24 , 19.70]	•
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega	26 81 1.70, df = 2 (P Z = 3.79 (P = balin (contro	119 402 9 = 0.43); 1 0.0001) biled relea	3 24 2 ² = 0%	59 273	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19]	
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega French 2014 Subtotal (99% CI) Total events:	26 81 1.70, df = 2 (P Z = 3.79 (P = balin (contro 4 4	119 402 9 = 0.43); 1 0.0001) blled relea 101	3 24 2 ² = 0% se) 2	59 273 110	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19] 2.18 [0.24 , 19.70]	
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega French 2014 Subtotal (99% CI)	26 81 1.70, df = 2 (P Z = 3.79 (P = b alin (contro 4 4 blicable	119 402 9 = 0.43); 1 0.0001) 0lled relea 101 101	3 24 2 ² = 0% se) 2	59 273 110	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19] 2.18 [0.24 , 19.70]	
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not app Test for overall effect: 1	26 81 1.70, df = 2 (P Z = 3.79 (P = balin (contro 4 4 blicable Z = 0.91 (P =	119 402 9 = 0.43); 1 0.0001) 0.0001) 0.0001 0.0001) 0.0001) 0.0001) 0.0001)	3 24 2 ² = 0% se) 2 2	59 273 110	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19] 2.18 [0.24 , 19.70]	
Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 1 Test for overall effect: 1 2.12.6 165 mg/d prega French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not app	26 81 1.70, df = 2 (P Z = 3.79 (P = balin (contro 4 4 blicable Z = 0.91 (P =	119 402 9 = 0.43); 1 0.0001) 0.0001) 0.0001 0.0001) 0.0001) 0.0001) 0.0001)	3 24 2 ² = 0% se) 2 2	59 273 110	14.7% 100.0%	4.30 [0.94 , 19.57] 2.35 [1.31 , 4.19] 2.18 [0.24 , 19.70]	

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 2.12. (Continued)

11CHCH 2014	U	114	4	110	100.070	2.09 [0.30 , 23.03	-	-			
Subtotal (99% CI)		114		110	100.0%	2.89 [0.36 , 23.05	5]	-			
Total events:	6		2								
Heterogeneity: Not applica	ble										
Test for overall effect: Z =	1.32 (P = 0.1	.9)									
							0.01	0.1	1	10	100
							Favours pr	egabalin	F	avours p	lacebo

Analysis 2.13. Comparison 2: Pregabalin versus placebo - subgroup analysis, Outcome 13: Weight gain

	Tiegau	alin	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
2.13.1 50 mg/d pregaba	alin						
French 2003	1	88	0	100	100.0%	3.40 [0.05 , 224.69]	
Subtotal (99% CI)		88		100	100.0%	3.40 [0.05 , 224.69]	
Total events:	1		0				
Heterogeneity: Not appl							
Test for overall effect: Z		0.45)					
2.13.2 150 mg/d pregal	palin						
Arroyo 2004	7	99	2	97	81.2%	3.43 [0.45 , 26.17]	
French 2003	2	88	0	100	18.8%	5.67 [0.11 , 301.50]	
Subtotal (99% CI)	_	187	-	197	100.0%	3.85 [0.64 , 23.35]	
Total events:	9	107	2	107	100.0 /0	5.65 [0.64], 25.55]	
Heterogeneity: Chi ² = 0		- 0 77). I					
Test for overall effect: Z			- 070				
2.13.3 300 mg/d pregal	nalin						
French 2003	6 G	90	0	100	100.0%	14.43 [0.34 , 620.87]	
Subtotal (99% CI)	0	90 90	0	100 100	100.0%	14.43 [0.34 , 620.87] 14.43 [0.34 , 620.87]	
Total events:	6	50	0	100	100.0 70	14.40 [0.04 , 020.07]	
Heterogeneity: Not appl			0				
Test for overall effect: Z		0.07)					
2.13.4 600 mg/d pregal	alin						
2.13.4 000 mg/u prega Arroyo 2004	Jann 13	92	2	97	16.7%	6.85 [1.00 , 46.76]	
Beydoun 2005	38	92 215	2	97 98	23.5%		—
•						8.66 [1.37 , 54.65]	
Elger 2005 Eronah 2002	28	137	5	73 100	55.8%	2.98 [0.90 , 9.84]	├──
French 2003	11	89 533	0	100	4.0%	25.81 [0.64 , 1046.44]	+
Subtotal (99% CI)	00	533	<i>c</i>	368	100.0%	5.88 [2.52 , 13.73]	
Total events:	90	0.05	9				
Heterogeneity: $Chi^2 = 3$	-		- = 15%				
Test for overall effect: Z	L = 5.39 (P < 1)						
			0 / 1				
2.13.5 Titrated dose of	pregabalin ((150 to 60	U .	<i>,</i>	10.00/		
2.13.5 Titrated dose of Baulac 2010	pregabalin 14	(150 to 60 152	2	141	18.6%	6.49 [0.95 , 44.46]	_ _
Test for overall effect: Z 2.13.5 Titrated dose of Baulac 2010 Elger 2005	pregabalin 14 25	(150 to 60 152 131	2 5	141 73	57.5%	2.79 [0.84 , 9.29]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009	pregabalin 14	(150 to 60 152 131 119	2	141 73 59	57.5% 23.9%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI)	pregabalin 14 25 14	(150 to 60 152 131	2 5 2	141 73	57.5%	2.79 [0.84 , 9.29]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events:	pregabalin 14 25 14 53	(150 to 60) 152 131 119 402	2 5 2 9	141 73 59	57.5% 23.9%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28]	•
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0	pregabalin (14 25 14 53 93, df = 2 (P	(150 to 60 152 131 119 402 = 0.63); I	2 5 2 9	141 73 59	57.5% 23.9%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28]	•
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0	pregabalin (14 25 14 53 93, df = 2 (P	(150 to 60 152 131 119 402 = 0.63); I	2 5 2 9	141 73 59	57.5% 23.9%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal	pregabalin (14 25 14 53 .93, df = 2 (P 2 = 3.74 (P =	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea	2^{2} 2^{5} 2^{2} $2^{2} = 0\%$ se)	141 73 59 273	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014	pregabalin (14 25 14 53 .93, df = 2 (P 53 .93, df = 2 (P	(150 to 60 152 131 119 402 = 0.63); I 0.0002)	2^{2} 5^{2} 2^{2} $9^{2} = 0\%$	141 73 59	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014	pregabalin (14 25 14 53 .93, df = 2 (P 2 = 3.74 (P =	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea	2^{2} 2^{5} 2^{2} $2^{2} = 0\%$ se)	141 73 59 273	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z	pregabalin (14 25 14 53 .93, df = 2 (P 2 = 3.74 (P =	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101	2^{2} 2^{5} 2^{2} $2^{2} = 0\%$ se)	141 73 59 273 110	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI) Total events:	pregabalin (14 25 14 53 93, df = 2 (P ; = 3.74 (P = balin (contro 1 1	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101	$2^{2} = 0^{2}$	141 73 59 273 110	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl	pregabalin (14 25 14 53 93, df = 2 (P = 3.74 (P = balin (contro 1 1 icable	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101 101	$2^{2} = 0^{2}$	141 73 59 273 110	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI)	pregabalin (14 25 14 53 93, df = 2 (P = 3.74 (P = palin (contro 1 icable = 0.73 (P =	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101 101 0.47)	$2^{2} = 0\%$ (1) (2) (2) (2) (2) (2) (2) (2) (2) (2) (2	141 73 59 273 110	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z 2.13.7 330 mg/d pregal	pregabalin (14 25 14 53 93, df = 2 (P = 3.74 (P = palin (contro 1 icable = 0.73 (P =	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101 101 0.47)	$2^{2} = 0\%$ (1) (2) (2) (2) (2) (2) (2) (2) (2) (2) (2	141 73 59 273 110	57.5% 23.9% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z 2.13.7 330 mg/d pregal French 2014	pregabalin (14 25 14 53 .93, df = 2 (P = 3.74 (P = balin (contro 1 icable = 0.73 (P = balin (contro	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101 101 0.47) Iled relea	$2^{2} = 0\%$ (2) (2) (2) (2) (2) (2) (2) (2) (2) (2)	141 73 59 273 110 110	57.5% 23.9% 100.0% 100.0% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86] 3.26 [0.05 , 215.86]	
2.13.5 Titrated dose of Baulac 2010 Elger 2005 Lee 2009 Subtotal (99% CI) Total events: Heterogeneity: Chi ² = 0 Test for overall effect: Z 2.13.6 165 mg/d pregal French 2014 Subtotal (99% CI) Total events: Heterogeneity: Not appl Test for overall effect: Z	pregabalin (14 25 14 53 .93, df = 2 (P = 3.74 (P = balin (contro 1 icable = 0.73 (P = balin (contro	(150 to 60 152 131 119 402 = 0.63); I 0.0002) Iled relea 101 101 0.47) Iled relea 114	$2^{2} = 0\%$ (2) (2) (2) (2) (2) (2) (2) (2) (2) (2)	141 73 59 273 110 110 110	57.5% 23.9% 100.0% 100.0% 100.0%	2.79 [0.84 , 9.29] 3.47 [0.52 , 23.28] 3.64 [1.49 , 8.87] 3.26 [0.05 , 215.86] 3.26 [0.05 , 215.86] 3.26 [0.05 , 215.86]	

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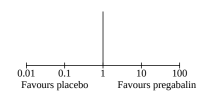
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Analysis 2.13. (Continued)

Heterogeneity: Not applicable Test for overall effect: Z = 1.84 (P = 0.07)



Comparison 3. Pregabalin versus active comparator

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3.1 50% or greater reduction in seizure frequency	3	1286	Risk Ratio (IV, Random, 95% CI)	1.03 [0.85, 1.25]
3.2 50% or greater reduction in seizure frequency - best-case analysis	3	1286	Risk Ratio (IV, Random, 95% CI)	1.60 [1.17, 2.19]
3.3 50% or greater reduction in seizure frequency - worst-case analysis	3	1286	Risk Ratio (M-H, Fixed, 95% CI)	0.67 [0.62, 0.74]
3.4 Seizure freedom	2	802	Risk Ratio (M-H, Fixed, 95% CI)	0.59 [0.37, 0.95]
3.5 Treatment withdrawal for any reason	3	1286	Risk Ratio (M-H, Fixed, 95% CI)	0.93 [0.76, 1.13]
3.6 Treatment withdrawal due to adverse effects	3	1286	Risk Ratio (M-H, Fixed, 95% CI)	1.04 [0.73, 1.48]
3.7 Ataxia	1		Risk Ratio (M-H, Fixed, 99% CI)	Totals not selected
3.8 Dizziness	3	1286	Risk Ratio (IV, Random, 99% CI)	1.64 [0.85, 3.16]
3.9 Fatigue	1		Risk Ratio (M-H, Fixed, 99% CI)	Totals not selected
3.10 Headache	3	1286	Risk Ratio (IV, Random, 99% CI)	0.83 [0.41, 1.65]
3.11 Nausea	1		Risk Ratio (M-H, Fixed, 99% CI)	Totals not selected
3.12 Somnolence	3	1286	Risk Ratio (M-H, Fixed, 99% CI)	1.16 [0.88, 1.53]
3.13 Weight gain	3	1286	Risk Ratio (IV, Random, 99% CI)	2.87 [0.94, 8.75]

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Analysis 3.1. Comparison 3: Pregabalin versus active comparator, **Outcome 1: 50% or greater reduction in seizure frequency**

	Pregal	balin	Active con	iparator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Baulac 2010	54	152	34	141	18.7%	1.47 [1.03 , 2.12]	
French 2016	134	242	140	242	41.2%	0.96 [0.82 , 1.12]	
Zaccara 2014	130	254	139	255	40.1%	0.94 [0.80 , 1.11]	
Total (95% CI)		648		638	100.0%	1.03 [0.85 , 1.25]	
Total events:	318		313				T
Heterogeneity: Tau ² = 0	0.02; Chi ² = 5	5.19, df = 2	(P = 0.07); I	² = 61%			0.5 0.7 1 1.5 2
Test for overall effect:	Z = 0.30 (P =	0.77)				Favours ac	tive comparator Favours pregaba

Test for subgroup differences: Not applicable

Analysis 3.2. Comparison 3: Pregabalin versus active comparator, Outcome 2: 50% or greater reduction in seizure frequency - best-case analysis

	Pregat	oalin	Active com	parator		Risk Ratio	Risk Ratio	,
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95	% CI
Baulac 2010	100	152	34	141	27.7%	2.73 [1.99 , 3.74]	-	
French 2016	188	242	140	242	36.4%	1.34 [1.18 , 1.52]		
Zaccara 2014	176	254	139	255	35.9%	1.27 [1.11 , 1.46]	-	
Total (95% CI)		648		638	100.0%	1.60 [1.17 , 2.19]		
Total events:	464		313				•	
Heterogeneity: Tau ² = 0.	.06; Chi ² = 1	9.54, df = 2	2 (P < 0.0001); I ² = 90%		0.01	0.1 1	10 100
Test for overall effect: Z	z = 2.98 (P =	0.003)				Favours active	comparator Fa	avours pregabalin
Test for subgroup different	ences: Not aj	pplicable						

Analysis 3.3. Comparison 3: Pregabalin versus active comparator, Outcome 3: 50% or greater reduction in seizure frequency - worst-case analysis

	Pregal	balin	Active com	parator		Risk Ratio	Risk l	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixe	d, 95% CI
Baulac 2010	54	152	74	141	16.4%	0.68 [0.52 , 0.88]		
French 2016	134	242	209	242	44.5%	0.64 [0.57 , 0.73]		
Zaccara 2014	130	254	184	255	39.1%	0.71 [0.62 , 0.82]		
Total (95% CI)		648		638	100.0%	0.67 [0.62 , 0.74]		
Total events:	318		467				•	
Heterogeneity: Chi ² = 1	.12, df = 2 (I	P = 0.57); I	$^{2} = 0\%$				0.5 0.7 1	1.5 2
Test for overall effect: 2	Z = 8.63 (P <	0.00001)				Favours ac	ctive comparator	Favours pregabalin
Test for subgroup differ	rences: Not a	pplicable						

Analysis 3.4. Comparison 3: Pregabalin versus active comparator, Outcome 4: Seizure freedom

	Pregal	balin	Active com	parator		Risk Ratio	Risk F	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed	l, 95% CI
Baulac 2010	6	152	4	141	9.9%	1.39 [0.40 , 4.83]		
Zaccara 2014	19	254	38	255	90.1%	0.50 [0.30 , 0.85]		
Total (95% CI)		406		396	100.0%	0.59 [0.37 , 0.95]		
Total events:	25		42				•	
Heterogeneity: Chi ² = 2	2.19, df = 1 (H	P = 0.14); I	² = 54%			0	.01 0.1 1	10 100
Test for overall effect: 2	Z = 2.19 (P =	0.03)				Favours ac	tive comparator	Favours pregabalin
Test for subgroup differ	rences: Not a	pplicable						

Analysis 3.5. Comparison 3: Pregabalin versus active comparator, Outcome 5: Treatment withdrawal for any reason

	Pregat	oalin	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Baulac 2010	46	152	40	141	26.7%	1.07 [0.75 , 1.52]	
French 2016	54	242	69	242	44.4%	0.78 [0.57 , 1.07]	_
Zaccara 2014	46	254	45	255	28.9%	1.03 [0.71 , 1.49]	
Total (95% CI)		648		638	100.0%	0.93 [0.76 , 1.13]	
Total events:	146		154				
Heterogeneity: Chi ² = 2	.04, df = 2 (F	9 = 0.36); I	2 = 2%				0.5 0.7 1 1.5 2
Test for overall effect: 2	Z = 0.73 (P =	0.46)				F	Favours pregabalin Favours active compara
Test for subgroup differ	ences: Not aj	pplicable					

Analysis 3.6. Comparison 3: Pregabalin versus active comparator, Outcome 6: Treatment withdrawal due to adverse effects

	Pregal	balin	Active com	parator		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
Baulac 2010	24	152	25	141	47.2%	0.89 [0.53 , 1.48]		
French 2016	16	242	15	242	27.3%	1.07 [0.54 , 2.11]		
Zaccara 2014	18	254	14	255	25.4%	1.29 [0.66 , 2.54]		
Total (95% CI)		648		638	100.0%	1.04 [0.73 , 1.48]		
Total events:	58		54				T	
Heterogeneity: Chi ² = 0	.75, df = 2 (I	P = 0.69); I	$^{2} = 0\%$			0.01	0.1 1 10	100
Test for overall effect: Z	Z = 0.22 (P =	0.82)				Favour	rs pregabalin Favours a	ctive comparator
Test for subgroup differ	ences: Not aj	pplicable						

Analysis 3.7. Comparison 3: Pregabalin versus active comparator, Outcome 7: Ataxia

Study or Subgroup	Pregal Events	oalin Total	Active Com Events	parator Total	Risk Ratio M-H, Fixed, 99% CI	Risk Ratio M-H, Fixed, 99% CI	
Baulac 2010	13	152	7	141	1.72 [0.54 , 5.55]	-+	
						0.001 0.1 1 10 avours pregabalin Favours	1000 active comparator

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	Pregal	balin	Active com	parator		Risk Ratio	Risk R	atio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 99% CI	IV, Random	, 99% CI
Baulac 2010	38	152	12	141	29.3%	2.94 [1.32 , 6.52]	-	
French 2016	22	242	20	242	30.5%	1.10 [0.51 , 2.35]		_
Zaccara 2014	56	254	39	255	40.2%	1.44 [0.89 , 2.34]		F
Total (99% CI)		648		638	100.0%	1.64 [0.85 , 3.16]		
Total events:	116		71					
Heterogeneity: Tau ² = 0	0.13; Chi ² = 5	.75, df = 2	(P = 0.06); I	2 = 65%		⊢ 0.0	1 0.1 1	10 100
Test for overall effect: 2	Z = 1.92 (P =	0.05)				Favo	urs pregabalin	Favours active comp

Analysis 3.8. Comparison 3: Pregabalin versus active comparator, Outcome 8: Dizziness

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Test for subgroup differences: Not applicable

Analysis 3.9. Comparison 3: Pregabalin versus active comparator, Outcome 9: Fatigue

Study or Subgroup	Pregal	balin	Active com	parator	Risk Ratio	Risk	Ratio
	Events	Total	Events	Total	M-H, Fixed, 99% CI	M-H, Fixe	d, 99% CI
Baulac 2010	26	152	14	141	. [,]	0.01 0.1 Savours pregabalin	i 10 100 Favours active comparator

Analysis 3.10. Comparison 3: Pregabalin versus active comparator, Outcome 10: Headache

	Pregal	balin	Active com	parator		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 99% CI	IV, Random, 99% Cl	
Baulac 2010	18	152	32	141	34.3%	0.52 [0.26 , 1.05]		
French 2016	17	242	20	242	30.4%	0.85 [0.38 , 1.92]		
Zaccara 2014	30	254	24	255	35.3%	1.25 [0.64 , 2.45]		
Total (99% CI)		648		638	100.0%	0.83 [0.41 , 1.65]		
Total events:	65		76					
Heterogeneity: Tau ² = 0	.14; Chi ² = 5	.50, df = 2	(P = 0.06); I ²	2 = 64%		0.01	0.1 1 10	100
Test for overall effect: Z	Z = 0.71 (P =	0.47)					s pregabalin Favours	s active comparato
Test for subgroup differ	ences: Not a	pplicable						

Analysis 3.11. Comparison 3: Pregabalin versus active comparator, Outcome 11: Nausea

	Prega	balin	Active com	parator	Risk Ratio	Risk F	Ratio
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 99% CI	M-H, Fixed	l, 99% CI
Zaccara 2014	3	254	15	255	0.20 [0.04 , 1.01]		
						0.01 0.1 1	
					F	avours pregabalin	Favours active comparator

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Analysis 3.12. Comparison 3: Pregabalin versus active comparator, Outcome 12: Somnolence

	Pregal	balin	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
Baulac 2010	30	152	15	141	12.7%	1.86 [0.87 , 3.95]	
French 2016	34	242	34	242	27.8%	1.00 [0.56 , 1.78]	_ _
Zaccara 2014	79	254	73	255	59.5%	1.09 [0.77 , 1.54]	•
Total (99% CI)		648		638	100.0%	1.16 [0.88 , 1.53]	
Total events:	143		122				T
Heterogeneity: Chi ² = 3	8.22, df = 2 (F	P = 0.20); I	² = 38%			0.01	
Test for overall effect: 2	Z = 1.37 (P =	0.17)					rs pregabalin Favours active comparate
Test for subgroup differ	rences: Not aj	pplicable					

Analysis 3.13. Comparison 3: Pregabalin versus active comparator, Outcome 13: Weight gain

	Pregal	balin	Active com	parator		Risk Ratio	Risk l	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 99% CI	IV, Randor	n, 99% CI
Baulac 2010	14	152	3	141	25.8%	4.33 [0.86 , 21.68]	_	
French 2016	19	242	13	242	41.1%	1.46 [0.60 , 3.58]	_	-
Zaccara 2014	24	254	5	255	33.0%	4.82 [1.39 , 16.74]		
Total (99% CI)		648		638	100.0%	2.87 [0.94 , 8.75]	-	
Total events:	57		21					•
Heterogeneity: Tau ² = 0).33; Chi ² = 4	.99, df = 2	(P = 0.08); I ²	$^{2} = 60\%$		0	0.01 0.1 1	10 100
Test for overall effect: 2	Z = 2.43 (P =	0.01)					vours pregabalin	Favours active comparat
Test for subgroup differ	rences: Not a	pplicable						

Comparison 4. Pregabalin versus active comparator - subgroup analysis

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.1 50% or greater reduction in seizure frequency	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.1.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	1.47 [1.03, 2.12]
4.1.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	0.94 [0.80, 1.11]
4.1.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.82, 1.12]
4.2 50% or greater reduction in seizure frequency - best-case analysis	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.2.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	2.73 [1.99, 3.74]
4.2.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	1.27 [1.11, 1.46]
4.2.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 95% Cl)	1.34 [1.18, 1.52]

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.3 50% or greater reduction in seizure frequency - worst-case analysis	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.3.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	0.68 [0.52, 0.88]
4.3.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	0.71 [0.62, 0.82]
4.3.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 95% CI)	0.64 [0.57, 0.73]
4.4 Seizure freedom	2		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.4.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	1.39 [0.40, 4.83]
4.4.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	0.50 [0.30, 0.85]
4.5 Treatment withdrawal for any reason	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.5.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	1.07 [0.75, 1.52]
4.5.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	1.03 [0.71, 1.49]
4.5.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 95% CI)	0.78 [0.57, 1.07]
4.6 Treatment withdrawal due to adverse effects	3		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.6.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 95% CI)	0.89 [0.53, 1.48]
4.6.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 95% CI)	1.29 [0.66, 2.54]
4.6.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 95% Cl)	1.07 [0.54, 2.11]
4.7 Dizziness	3		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
4.7.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 99% CI)	2.94 [1.32, 6.52]
4.7.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 99% CI)	1.44 [0.89, 2.34]
4.7.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 99% Cl)	1.10 [0.51, 2.35]
4.8 Headache	3		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only

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Cochrane Database of Systematic Reviews

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
4.8.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 99% CI)	0.52 [0.26, 1.05]
4.8.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 99% CI)	1.25 [0.64, 2.45]
4.8.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 99% CI)	0.85 [0.38, 1.92]
4.9 Nausea	1		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
4.9.1 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 99% CI)	0.20 [0.04, 1.01]
4.10 Somnolence	3		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
4.10.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 99% CI)	1.99 [0.91, 4.33]
4.10.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 99% CI)	1.09 [0.77, 1.54]
4.10.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 99% CI)	1.00 [0.56, 1.78]
4.11 Weight gain	3		Risk Ratio (M-H, Fixed, 99% CI)	Subtotals only
4.11.1 Pregabalin versus lamotrig- ine	1	293	Risk Ratio (M-H, Fixed, 99% CI)	4.33 [0.86, 21.68]
4.11.2 Pregabalin versus levetirac- etam	1	509	Risk Ratio (M-H, Fixed, 99% CI)	4.82 [1.39, 16.74]
4.11.3 Pregabalin versus gabapentin	1	484	Risk Ratio (M-H, Fixed, 99% CI)	1.46 [0.60, 3.58]



Analysis 4.1. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 1: 50% or greater reduction in seizure frequency

	Pregab	alin	Active com	parator		Risk Ratio	Risk R	atio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed	, 95% CI
4.1.1 Pregabalin versu	s lamotrigine	2						
Baulac 2010	54	152	34	141	100.0%	1.47 [1.03 , 2.12]		
Subtotal (95% CI)		152		141	100.0%	1.47 [1.03 , 2.12]		
Total events:	54		34					•
Heterogeneity: Not appl	licable							
Test for overall effect: Z	L = 2.09 (P = 0)	0.04)						
4.1.2 Pregabalin versu	s levetiraceta	ım						
Zaccara 2014	130	254	139	255	100.0%	0.94 [0.80 , 1.11]		
Subtotal (95% CI)		254		255	100.0%	0.94 [0.80 , 1.11]	•	I
Total events:	130		139					
Heterogeneity: Not appl	licable							
Test for overall effect: Z	L = 0.75 (P = 0.75)	0.45)						
4.1.3 Pregabalin versu	s gabapentin							
French 2016	134	242	140	242	100.0%	0.96 [0.82 , 1.12]		
Subtotal (95% CI)		242		242	100.0%	0.96 [0.82 , 1.12]		
Total events:	134		140				The second se	
Heterogeneity: Not appl	licable							
Test for overall effect: Z	L = 0.55 (P = 0.55)	0.58)						
Test for subgroup differ	ences: Chi² =	5.19, df =	2 (P = 0.07)	, I ² = 61.4%)	0.01 Favours activ	L 0.1 1 ve comparator	10 100 Favours pregabalii

Analysis 4.2. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 2: 50% or greater reduction in seizure frequency - best-case analysis

	Favours active co	mparator	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
4.2.1 Pregabalin versus la	amotrigine						
Baulac 2010	100	152	34	141	100.0%	2.73 [1.99 , 3.74]	
Subtotal (95% CI)		152		141	100.0%	2.73 [1.99 , 3.74]	
Total events:	100		34				•
Heterogeneity: Not applica	able						
Test for overall effect: $Z =$	6.26 (P < 0.00001)						
4.2.2 Pregabalin versus le	evetiracetam						
Zaccara 2014	176	254	139	255	100.0%	1.27 [1.11 , 1.46]	•
Subtotal (95% CI)		254		255	100.0%	1.27 [1.11 , 1.46]	•
Total events:	176		139				¥
Heterogeneity: Not applica	able						
Test for overall effect: Z =	3.39 (P = 0.0007)						
4.2.3 Pregabalin versus ga	abapentin						
French 2016	188	242	140	242	100.0%	1.34 [1.18 , 1.52]	
Subtotal (95% CI)		242		242	100.0%	1.34 [1.18 , 1.52]	A
Total events:	188		140				•
Heterogeneity: Not applica	able						
Test for overall effect: Z =	4.55 (P < 0.00001)						
Test for subgroup difference	ces: Chi ² = 19.54, d	f = 2 (P < 0.00)	01), I ² = 89.8	%		0.01	0.1 1 10 10
						Favours activ	e comparator Favours pregab

Pregabalin add-on for drug-resistant focal epilepsy (Review)



Analysis 4.3. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 3: 50% or greater reduction in seizure frequency - worst-case analysis

	Pregab	alin	Active com	parator		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% C	I
4.3.1 Pregabalin versu	s lamotrigine	2						
Baulac 2010	54	152	74	141	100.0%	0.68 [0.52 , 0.88]		
Subtotal (95% CI)		152		141	100.0%	0.68 [0.52 , 0.88]		
Total events:	54		74				•	
Heterogeneity: Not app	licable							
Test for overall effect: Z	Z = 2.88 (P = 0)	0.004)						
4.3.2 Pregabalin versu	ıs levetiraceta	ım						
Zaccara 2014	130	254	184	255	100.0%	0.71 [0.62 , 0.82]		
Subtotal (95% CI)		254		255	100.0%	0.71 [0.62 , 0.82]	•	
Total events:	130		184				•	
Heterogeneity: Not app	licable							
Test for overall effect: Z	Z = 4.73 (P < 0	0.00001)						
4.3.3 Pregabalin versu	ıs gabapentin							
French 2016	134	242	209	242	100.0%	0.64 [0.57 , 0.73]		
Subtotal (95% CI)		242		242	100.0%	0.64 [0.57 , 0.73]	•	
Total events:	134		209				•	
Heterogeneity: Not app	licable							
Test for overall effect: Z	Z = 7.04 (P < 0)	0.00001)						
Test for subgroup differ	ences: Chi ² –	1 11 df -	2 (D - 0 57)	12 - 0%		, H		
rest for subgroup differ	ences, Gill [®] -	1.11, ul –	2(r - 0.37)	,1 - 070		0.01 Favours activ) 100 s pregabal
						Favours acus	ravoui	s pregaua

Analysis 4.4. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 4: Seizure freedom

	Pregab	alin	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
4.4.1 Pregabalin versus	lamotrigin	2					
Baulac 2010	6	152	4	141	100.0%	1.39 [0.40 , 4.83]	
Subtotal (95% CI)		152		141	100.0%	1.39 [0.40 , 4.83]	
Total events:	6		4				
Heterogeneity: Not appli	cable						
Test for overall effect: Z	= 0.52 (P =	0.60)					
4.4.2 Pregabalin versus	levetiraceta	ım					
Zaccara 2014	19	254	38	255	100.0%	0.50 [0.30 , 0.85]	
Subtotal (95% CI)		254		255	100.0%	0.50 [0.30 , 0.85]	
	19		38				•
Total events:	19		50				
			50				
Total events: Heterogeneity: Not appli Test for overall effect: Z	cable	0.010)	50				



Analysis 4.5. Comparison 4: Pregabalin versus active comparator subgroup analysis, Outcome 5: Treatment withdrawal for any reason

Study or Subgroup	Prega Events	balin Total	Active com Events	iparator Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk Ratio M-H, Fixed, 95% CI
4.5.1 Pregabalin versu	ıs lamotrigir	1e					
Baulac 2010	46	152	40	141	100.0%	1.07 [0.75 , 1.52]	
Subtotal (95% CI)		152		141	100.0%	1.07 [0.75 , 1.52]	
Total events:	46		40				Ť
Heterogeneity: Not app	licable						
Test for overall effect: 2	Z = 0.36 (P =	0.72)					
4.5.2 Pregabalin versu	ıs levetirace	tam					
Zaccara 2014	46	254	45	255	100.0%	1.03 [0.71 , 1.49]	
Subtotal (95% CI)		254		255	100.0%	1.03 [0.71 , 1.49]	
Total events:	46		45				Ť
Heterogeneity: Not app	licable						
Test for overall effect:	Z = 0.14 (P =	• 0.89)					
4.5.3 Pregabalin versu	ıs gabapenti	n					
French 2016	54	242	69	242	100.0%	0.78 [0.57 , 1.07]	
Subtotal (95% CI)		242		242	100.0%	0.78 [0.57 , 1.07]	
Total events:	54		69				•
Heterogeneity: Not app	licable						
Test for overall effect: 2		• 0.12)					
Test for subgroup differ	rences: Chi²	= 2.04, df =	= 2 (P = 0.36)	, I ² = 2.0%		I	0.01 0.1 1 10 100 Favours pregabalin Favours active compa

Analysis 4.6. Comparison 4: Pregabalin versus active comparator subgroup analysis, Outcome 6: Treatment withdrawal due to adverse effects

	Prega	balin	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
4.6.1 Pregabalin versu	s lamotrigir	ie					
Baulac 2010	24	152	25	141	100.0%	0.89 [0.53 , 1.48]	-
Subtotal (95% CI)		152		141	100.0%	0.89 [0.53 , 1.48]	
Total events:	24		25				Ţ
Heterogeneity: Not appl	icable						
Test for overall effect: Z	L = 0.44 (P =	0.66)					
4.6.2 Pregabalin versu	s levetirace	tam					
Zaccara 2014	18	254	14	255	100.0%	1.29 [0.66 , 2.54]	
Subtotal (95% CI)		254		255	100.0%	1.29 [0.66 , 2.54]	
Total events:	18		14				
Heterogeneity: Not appl	icable						
Test for overall effect: Z	L = 0.74 (P =	0.46)					
4.6.3 Pregabalin versu	s gabapenti	n					
French 2016	16	242	15	242	100.0%	1.07 [0.54 , 2.11]	-
Subtotal (95% CI)		242		242	100.0%	1.07 [0.54 , 2.11]	
Total events:	16		15				T
Heterogeneity: Not appl	icable						
Test for overall effect: Z	z = 0.19 (P =	0.85)					
		-					
Test for subgroup differ	ences: Chi ² :	= 0.75, df =	= 2 (P = 0.69)	$I^2 = 0\%$		C	0.01 0.1 1 10 100
0 1			. ,				vours pregabalin Favours active compa

Analysis 4.7. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 7: Dizziness

	Pregal	balin	Active com	parator		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
4.7.1 Pregabalin versus	lamotrigin	ie					
Baulac 2010	38	152	12	141	100.0%	2.94 [1.32 , 6.52]	
Subtotal (99% CI)		152		141	100.0%	2.94 [1.32 , 6.52]	
Total events:	38		12				•
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 3.48 (P =	0.0005)					
4.7.2 Pregabalin versus	levetiracet	am					
Zaccara 2014	56	254	39	255	100.0%	1.44 [0.89 , 2.34]	-
Subtotal (99% CI)		254		255	100.0%	1.44 [0.89 , 2.34]	▲
Total events:	56		39				•
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 1.94 (P =	0.05)					
4.7.3 Pregabalin versus	gabapenti	n					
French 2016	22	242	20	242	100.0%	1.10 [0.51 , 2.35]	
Subtotal (99% CI)		242		242	100.0%	1.10 [0.51 , 2.35]	
Total events:	22		20				Ť
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 0.32 (P =	0.75)					
Test for subgroup differer	ıces: Chi² =	= 5.75. df =	= 2 (P = 0.06).	. I² = 65.2%		0.	
and an area		5 5 , ui	- (- 0100)	, - 001270	-	••	purs pregabalin Favours active comp

Analysis 4.8. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 8: Headache

	Pregal		Active com	-		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
4.8.1 Pregabalin versus l	lamotrigin	e					
Baulac 2010	18	152	32	141	100.0%	0.52 [0.26 , 1.05]	
Subtotal (99% CI)		152		141	100.0%	0.52 [0.26 , 1.05]	—
Total events:	18		32				•
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 2.41 (P =	0.02)					
4.8.2 Pregabalin versus l	levetiracet	am					
Zaccara 2014	30	254	24	255	100.0%	1.25 [0.64 , 2.45]	
Subtotal (99% CI)		254		255	100.0%	1.25 [0.64 , 2.45]	
Total events:	30		24				
Heterogeneity: Not applic	able						
Test for overall effect: Z =	= 0.88 (P =	0.38)					
4.8.3 Pregabalin versus g	gabapentiı	1					
French 2016	17	242	20	242	100.0%	0.85 [0.38 , 1.92]	
Subtotal (99% CI)		242		242	100.0%	0.85 [0.38 , 1.92]	
Total events:	17		20				•
Heterogeneity: Not applic	able						
	= 0.51 (P =	0.01)					

Analysis 4.9. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 9: Nausea

Pregal	balin	Active com	parator		Risk Ratio	Risk Ratio	
Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99%	6 CI
levetiracet	am						
3	254	15	255	100.0%	0.20 [0.04 , 1.01]		
	254		255	100.0%	0.20 [0.04 , 1.01]		
3		15					
cable							
= 2.56 (P =	0.01)						
nces: Not a	pplicable				0.0	01 0.1 1	10 100
					Fave	ours pregabalin Fa	vours active compara
	Events levetiracet 3 cable = 2.56 (P =	levetiracetam 3 254 254 3	Events Total Events levetiracetam 3 254 15 254 3 15 3 15 15 cable = 2.56 (P = 0.01) 15	Events Total Events Total levetiracetam 3 254 15 255 254 254 255 3 15 cable = 2.56 (P = 0.01) 15 16	Events Total Events Total Weight levetiracetam 3 254 15 255 100.0% 254 254 255 100.0% 3 15 cable = 2.56 (P = 0.01)	Events Total Events Total Weight M-H, Fixed, 99% CI levetiracetam 3 254 15 255 100.0% 0.20 [0.04, 1.01] 254 255 100.0% 0.20 [0.04, 1.01] 3 15 cable = 2.56 (P = 0.01) 0.01 0.01	Events Total Events Total Weight M-H, Fixed, 99% CI M-H, Fixed, 99% levetiracetam 3 254 15 255 100.0% 0.20 [0.04, 1.01]

Analysis 4.10. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 10: Somnolence

		Active com	parator		Risk Ratio	Risk Ratio
vents	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI
amotrigii	1e					
30	152	14	141	100.0%	1.99 [0.91 , 4.33]	+ - -
	152		141	100.0%	1.99 [0.91 , 4.33]	
30		14				•
ole						
2.28 (P =	0.02)					
evetirace	tam					
79	254	73	255	100.0%	1.09 [0.77 , 1.54]	•
	254		255	100.0%	1.09 [0.77 , 1.54]	—
79		73				T
ole						
).61 (P =	0.54)					
abapenti	n					
34	242	34	242	100.0%	1.00 [0.56 , 1.78]	
	242		242	100.0%	1.00 [0.56 , 1.78]	
34		34				Ť
ole						
).00 (P =	1.00)					
	30 30 1e .28 (P = 1 evetirace 79 79 1e .61 (P = 1 34 34 34	152 30 le .28 (P = 0.02) vetiracetam 79 254 254 79 le .61 (P = 0.54) abapentin 34 242 242 34 le	30 152 14 152 14 30 14 le 14 .28 (P = 0.02) 14 vettracetam 79 79 254 73 254 73 254 73 16 9 .54 .61 (P = 0.54) 34 242 34 242 34 34 242 34 34 34 34 le .34 34	30 152 14 141 30 14 141 30 14 14 abe 14 14 .28 (P = 0.02) 255 257 254 73 255 254 255 79 73 255 .61 (P = 0.54) 73 255 34 242 34 242 34 34 34 242 34 34 34 34 le 54 34 34	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

Analysis 4.11. Comparison 4: Pregabalin versus active comparator - subgroup analysis, Outcome 11: Weight gain

	Prega	balin	Active com	parator		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 99% CI	M-H, Fixed, 99% CI	
4.11.1 Pregabalin versu	s lamotrigi	ine						
Baulac 2010	14	152	3	141	100.0%	4.33 [0.86 , 21.68]	↓ ■	
Subtotal (99% CI)		152		141	100.0%	4.33 [0.86 , 21.68]		
Total events:	14		3					
Heterogeneity: Not appli	cable							
Test for overall effect: Z	= 2.34 (P =	0.02)						
4.11.2 Pregabalin versu	s levetiraco	etam						
Zaccara 2014	24	254	5	255	100.0%	4.82 [1.39 , 16.74]		
Subtotal (99% CI)		254		255	100.0%	4.82 [1.39 , 16.74]		
Total events:	24		5					
Heterogeneity: Not appli	cable							
Test for overall effect: Z	= 3.25 (P =	0.001)						
4.11.3 Pregabalin versu	s gabapent	tin						
French 2016	19	242	13	242	100.0%	1.46 [0.60 , 3.58]		
Subtotal (99% CI)		242		242	100.0%	1.46 [0.60 , 3.58]		
Total events:	19		13				—	
Heterogeneity: Not appli	cable							
	= 1.09 (P =	0.28)						

APPENDICES

Appendix 1. Cochrane Register of Studies (CRS Web) search strategy

1. MESH DESCRIPTOR Pregabalin EXPLODE ALL AND CENTRAL: TARGET

2. (lyrica OR pregabalin*):AB,KW,KY,MC,MH,TI AND CENTRAL:TARGET

3. #1 OR #2

- 4. MESH DESCRIPTOR Epilepsies, Partial EXPLODE ALL AND CENTRAL: TARGET
- 5. ((partial or focal) and (seizure* or epilep*)):AB,KW,KY,MC,MH,TI AND CENTRAL:TARGET

6. #4 OR #5

7. #3 AND #6

8. (monotherap* NOT (adjunct* OR "add-on" OR "add on" OR adjuvant* OR combination* OR polytherap*)):TI AND CENTRAL:TARGET

9. #7 NOT #8

10. #9 AND >05/07/2018:CRSCREATED

Appendix 2. MEDLINE (Ovid) search strategy

This strategy includes a modification of the Cochrane Highly Sensitive Search Strategy for identifying randomised trials (Lefebvre 2021).

- 1. exp Pregabalin/ or (pregabalin\$ or lyrica).tw.
- 2. exp Epilepsies, Partial/
- 3. ((partial or focal) and (seizure\$ or epilep\$)).tw.

4. 2 or 3

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- 5.1 and 4
- 6. exp controlled clinical trial/ or (randomi?ed or placebo or randomly).ab.
- 7. clinical trials as topic.sh.
- 8. trial.ti.
- 9.6 or 7 or 8
- 10. exp animals/ not humans.sh.
- 11. 9 not 10
- 12. 5 and 11
- 13. (monotherap\$ not (adjunct\$ or "add-on" or "add on" or adjuvant\$ or combination\$ or polytherap\$)).ti.
- 14. 12 not 13
- 15. limit 14 to ed=20180705-20201116
- 16. 14 not (1\$ or 2\$).ed.
- 17. 16 and (2018\$ or 2019\$ or 2020\$).dt.
- 18. 15 or 17
- 19. remove duplicates from 18

WHAT'S NEW

Date	Event	Description
1 April 2022	Amended	Minor typos corrected.

HISTORY

Protocol first published: Issue 1, 2006 Review first published: Issue 1, 2008

Date	Event	Description
16 November 2020	New search has been performed	Searches updated 16 November 2020; two new studies (Antinew 2019; Mann 2020) have been added to the review.
16 November 2020	New citation required but conclusions have not changed	Conclusions remain the same.
5 July 2018	New search has been performed	Searches updated 5 July 2018; three new studies (French 2014, French 2016 and Zaccara 2014) have been added to the review.
		The term 'partial' has been replaced by 'focal', in accordance with the most recent classification of epilepsies of the Interna- tional League Against Epilepsy (Scheffer 2017).
5 July 2018	New citation required but conclusions have not changed	Conclusions remain the same.

Pregabalin add-on for drug-resistant focal epilepsy (Review)

Date	Event	Description
9 January 2014	New search has been performed	Searches updated 9 January 2014; one previously ongoing study, IRCT2012091210508N4, has been added to excluded studies (Taghdiri 2015).
9 January 2014	New citation required but conclusions have not changed	Conclusions remain the same.
12 June 2012	New search has been performed	Two new studies were included in this update of the original re- view.
7 August 2009	Amended	Copy edits made at editorial base.
16 September 2008	Amended	Converted to new review format.

CONTRIBUTIONS OF AUTHORS

MP and RB carried out and completed the update of this review. MP and RB assessed trials for eligibility and completed data extraction. RB and MP both contributed to the writing of the review and the data analysis. AGM provided supervision throughout the review process.

DECLARATIONS OF INTEREST

MP: none known.

RB: none known.

AGM: a consortium of pharmaceutical companies (GSK, EISAI, UCB Pharma) funded the National Audit of Seizure Management in Hospitals (NASH) through grants paid to the University of Liverpool.

Professor Marson is funded in part by The National Institute for Health Research Collaboration for Leadership in Applied Health Research and Care North West Coast (NIHR CLAHRC NWC).

Professor Marson is a National Institute for Health Research (NIHR) Senior Investigator. The views expressed in this article are those of the author(s) and not necessarily those of the NIHR, or the Department of Health and Social Care.

Professor Marson is the Co-ordinating Editor of the Cochrane Epilepsy Group; however, he was not involved in the editorial process of this review update.

SOURCES OF SUPPORT

Internal sources

• No sources of support provided

External sources

• National Institute for Health Research (NIHR), UK

This review update was supported by the National Institute for Health Research, via Cochrane Infrastructure funding to the Epilepsy Group. The views and opinions expressed therein are those of the author(s) and do not necessarily reflect those of the Systematic Reviews Programme, NIHR, NHS or the Department of Health and Social Care.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The original protocol was amended for the previous review to include interventions comparing pregabalin to other antiepileptic drugs.

The method of analysis for examining dose regression was changed for the previous version of the review due to advances in techniques for analysis binary data. Specifically, a generalised linear mixed model using the software package STATA SE version 14 (Stata) was employed as opposed to a generalised linear model. We continued to use a generalised linear mixed model in the current review update.

The title of the review has been changed from "Pregabalin add-on for drug-resistant partial epilepsy" to "Pregabalin add-on for drugresistant focal epilepsy" in accordance with the latest classification of epilepsies released by the International League Against Epilepsy (ILAE) (Scheffer 2017). Likewise, any previous mention of "partial epilepsy" or "refractory epilepsy" throughout this review was changed to "focal epilepsy" and "drug-resistant epilepsy", respectively.



INDEX TERMS

Medical Subject Headings (MeSH)

*Drug Resistant Epilepsy [drug therapy]; Drug Therapy, Combination; *Epilepsies, Partial [drug therapy]; Pregabalin [therapeutic use]; Randomized Controlled Trials as Topic

MeSH check words

Humans