National Institute for Health and Care Excellence

Draft for consultation

Epilepsies in children, young people and adults

[G] Effectiveness of antiseizure therapies in the treatment of absence seizures

NICE guideline number tbc

Evidence reviews underpinning recommendations 5.3.1-5.3.9 in the NICE guideline

November 2021

Draft for Consultation

These evidence reviews were developed by the National Guideline Alliance which is a part of the Royal College of Obstetricians and Gynaecologists



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ISBN:

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Effectiveness of antiseizure therapies in the treatment of absence seizures

3 Review question

- 4 What antiseizure therapies (monotherapy or add-on) are effective in the treatment of ab-
- 5 sence seizures?

6 Introduction

- Absence seizures are a form of generalised epileptic seizure, characterised by behavioural
- 8 arrest with lack of awareness. Typical absences are sudden in onset, last a short duration of
- 9 time (5 to 10 seconds), and stop abruptly. They are associated with a characteristic electro-
- 10 encephalogram (EEG) pattern of regular 3 to 4 per second spike and wave activity. They are
- 11 characteristic of several epilepsy syndromes: in children, childhood absence epilepsy where
- they occur multiple times per day, and later onset juvenile absence epilepsy where they oc-
- cur less frequently. In adults they may be a feature of juvenile myoclonic epilepsy. They may
- 14 be seen in other epilepsy syndromes with other seizure types. Atypical absence seizures
- may be less obvious, last longer, and be associated with more variable patterns on EEG.
- They typically do not occur in isolation, but with other seizure types as part of more complex
- 17 epilepsies. The aim of this review is to determine which antiseizure therapies improve out-
- 18 comes in people with epilepsy who have absence seizures.

19 Summary of the protocol

- 20 Please see Table 1 for a summary of the Population, Intervention, Comparison and Outcome
- 21 (PICO) characteristics of this review.

22 Table 1: Summary of the protocol (PICO table)

| Population | People with confirmed epilepsy with absence seizures |
|--------------|--|
| | |
| Intervention | The following treatments and their combinations will be considered: • Acetazolamide • Clobazam • Clonazepam |
| | Ethosuximide |
| | Ketogenic diet (included as this is an accepted first or second line treatment for these type of seizures) |
| | Lamotrigine |
| | Levetiracetam |
| | Methsuximide/mesuximide |
| | Sodium Valproate |
| | Topiramate |
| | Zonisamide |
| Comparison | Any of the above and their combinationsNo treatment/placebo |
| Outcomes | Critical |
| | Seizure freedom (12 months data and short term, (minimum 3 months with 100% freedom) of starting treatment). |
| | 24 hour EEG seizure freedom |

- Time to withdrawal of treatment or change of medication (for example, because of uncontrollable seizures)
- Adverse events, as assessed by:
 - $_{\odot}$ % of patients with reported side effects (trial defined adverse and serious adverse effects)
 - treatment cessation due to adverse event (dichotomous outcome only)
 - o mortality

Important

- Neuropsychological changes (IQ testing, or other validated tools)
- Health-related quality of life (validated tools only)
- 1 EEG: electroencephalogram; IQ: intelligence quotient
- 2 For further details see the review protocol in appendix A.

3 Methods and process

- 4 This evidence review was developed using the methods and process described in <u>Develop-</u>
- 5 <u>ing NICE guidelines: the manual</u>. Methods specific to this review question are described in
- 6 the review protocol in appendix A and the methods document (supplementary document 1).
- 7 Declarations of interest were recorded according to <u>NICE's conflicts of interest policy.</u>

8 Clinical evidence

9 Included studies

- 10 Seven studies (reporting on six randomised controlled trials [RCTs]) were identified for inclu-
- sion in this review. Four studies reporting on 3 RCTs compared ethosuximide to sodium
- valproate (Callaghan 1982, Glauser 2010, Glauser 2013, Sato 1982); 3 studies reporting on
- 13 2 RCTs compared lamotrigine to sodium valproate (Coppola 2004, Glauser 2010, Glauser
- 14 2013); 1 RCT compared_levetiracetam to placebo (Fattore 2011); 1 RCT compared lamotrig-
- ine to placebo (Frank 1999) and 2 studies reportinging on 1 RCT compared ethosuximide to
- 16 lamotrigine (Glauser 2010, Glauser 2013).
- 17 The included studies are summarised in Table 2 to Table 6.

18

19 See the literature search strategy in appendix B and study selection flow chart in appendix C.

20 Excluded studies

- 21 Studies not included in this review with reasons for their exclusions are provided in appendix
- 22 K.

26

27

23 Summary of clinical studies included in the evidence review

Summaries of the studies that were included in this review are presented in Table 2 to Table 6.

Table 2: Summary of included studies. Comparison 1: ethosuximide versus sodium valproate

| | P | | | |
|-----------|----------------|--------------|------------------|------------------------|
| Study | Population | Intervention | Comparison | Outcomes |
| Callaghan | N = 28 peo- | Ethosuximide | Sodium valproate | Seizure remission |
| 1982 | ple with typi- | n=14 | n=14 | (complete, partial, no |

| Study | Population | Intervention | Comparison | Outcomes |
|---|--|---|---|--|
| RCT | cal absence seizures Mean age: ESM: 8 years (range 4 to14) VPA: 9 years (range 5 to 15) | Initial dose of 250mg/day Dose increased by 250mg/day to a maximum 1500mg/day as required | Initial dose of 400mg/day Dose increased by 200mg/day to a max- imum of 2400mg/day | remission) • EEG response - normal • Adverse events - any |
| Glauser 2010 Multi-centre RCT US Reports 20 week follow-up data | N = 453 people with absence sei- zures (three- arm trial) Mean age: not provided | Ethosuximide n=156 250mg capsules or 250 mg per 5 ml of syrup. Highest allowable daily dose of 60mg/kg (to a max- imum of 2000mg). | Sodium valproate n=148 25mg capsules or 125mg dose of sprinkles. Highest allowable daily dose of 60mg/kg (to a maximum of 3000mg). | Freedom from treatment failure Adverse events (serious adverse events; intolerable adverse effects) Study drug discontinued Withdrawal from study Attentional dysfunction (proxy outcome for neuropsychological changes) |
| Glauser 2013 See Glauser 2010 Reports 12 month follow-up data | See Glauser 2010 | See Glauser 2010 | See Glauser 2010 | Freedom from treatment failure Adverse events (serious adverse events; intolerable adverse effects) Attentional dysfunction (proxy outcome for neuropsychological changes) Withdrawal from study |
| Sato 1982 Cross-over RCT; only period 1 was reported and synthesised US | N = 45 people with absence seizures (treatment naive n=16; refractory n=29) First line therapy (naive) and addon therapy (refractory) Mean age: 11.7 years | Ethosuximide plus placebo n=23 (treatment naïve n=9, refractory n=14) Dosing schedule group 1: Daily dose of 250-1500mg Dosing schedule group 2: Daily dose of 250 to 1500mg Naive patients with 100% seizure control and refractory | Sodium valproate plus placebo n=22 (treatment na- ïve n=7, refractory n=15) Dosing schedule group 1: Initial dose of 15-20mg/kg for 5 days, increased to maximum of 30mg/kg if 12 hour telemetered EEG still showed generalised spike-wave discharges. | Seizure freedom |

2

| Study | Population | Intervention | Comparison | Outcomes |
|-------|-----------------|--|---|----------|
| | (range 4 to 18) | patients with at least 80% control during the first 6 weeks of treatment were not crossed over to alternative treatment because of ethical concerns regarding with-drawal of effective medications. Instead, they were maintained on the same drug for 3 months in a double-blind manner. Only those patients who did not 'respond' as described above; or experienced serious adverse reactions to the first therapy were crossed over to the alternative treatment. | Dosing schedule group 2: Initial dose of 12.5 to 20mg/kg, with a dosage increase every 2 days for 2 weeks up to a maximum daily dosage of 60mg/kg. Naive patients with 100% seizure control and refractory patients with at least 80% control during the first 6 weeks of treatment were not crossed over to alternative treatment because of ethical concerns regarding withdrawal of effective medications. Instead, they were maintained on the same drug for 3 months in a double-blind manner. Only those patients who did not 'respond' as described above; or experienced serious adverse reactions to the first therapy were crossed over to the alternative treatment. | |

EEG: electroencephalogram; ESM: ethosuximide; RCT: randomised controlled trial; VPA: sodium valproate

Table 3: Summary of included studies. Comparison 2: lamotrigine versus sodium valproate

| Study | Population | Intervention | Comparison | Outcomes |
|-----------------|---|---|--|--|
| Coppola 2004 | N = 38 peo- ple with ab- sence sei- | <u>Lamotrigine</u> n=19 | Sodium valproate n=19 | Seizure freedom (1, 3 and 12 months)Adverse events - any |
| RCT | zures Mean age: VPA: 7.5 years (range 3 to 13) LTG: 7.5 | Initial dose of 0.5mg/kg/day for 2 weeks, followed by 1.0mg/kg/day for 2 weeks. Doses then increased by 1mg/kg/day every 5 days until seizures were under control | Initial dose of 10/mg/kg/day and increased by 5mg/kg/day every 3 days until seizures were under control | Treatment cessation due to adverse events Withdrawal from study |
| | years (range 4 to 12) | were under control | | |

| Ottorile | Damulatian | Internetion | 0 | 0.45.5 |
|--|--|--|---|---|
| Study | Population | Intervention | Comparison | Outcomes |
| Glauser 2010 Multi-centre | N = 453 people with absence sei- | <u>Lamotrigine</u> n=149 | Sodium valproate n=148 | Freedom from treatment failureAdverse events (se- |
| RCT | zures (three- arm trial) | 5mg and 25mg chewable tablets. Maximum dose of | 25mg capsules or 125mg dose of sprinkles. | rious adverse events; intolerable adverse effects) |
| | Mean age: not provided | 12 mg/kg up to a maximum of | Highest allowable | Study drug discontinued |
| Reports 20 week follow- up data | · | 600mg/day | daily dose of 60mg/kg (to a maxi- | Withdrawal from study |
| | | | mum of 3000mg). | Attentional dysfunction (proxy outcome for neuropsychological changes) |
| Glauser 2013 | See Glauser 2010 | See Glauser 2010 | See Glauser 2010 | Freedom from treatment failure |
| See Glauser 2010 | | | | Adverse events (serious adverse events; intolerable adverse effects) |
| Reports 12 month follow- up data | | | | Attentional dysfunction (proxy outcome for neuropsychological changes) Withdrawal from |
| | | | | study |

¹ LTG: lamotrigine; RCT: randomised controlled trial; VPA: sodium valproate

2 Table 4: Summary of included studies. Comparison 3: levetiracetam versus placebo

| Table 4. Outliniary of included studies. Comparison 5. levelifacetain versus placeso | | | | | |
|--|--|---|--|---|--|
| Study | Population | Intervention | Comparison | Outcomes | |
| Fattore 2011 Multi-centre RCT Italy | N = 59 people with absence seizures Mean age LEV: 8.7 years (range 4.9 to 13) Placebo: 7.9 years (range 4 to 15) | Levetiracetam n=38 Initial dosage of 10mg/kg/day for 3 days followed by 15mg/kg/day for 4 days. If accepted and seizures continued dosage increased to 20mg/kg/day and maintained until day 14 | Placebo n=21 Equivalent tablet or liquid formulation given. | Responder status (free from clinical seizures on days 13 and 14; and free from EEG seizures on day 14) Patients free from clinical and EEG seizures (days 4 – 7; days 11 - 14) 50% reduction vs baseline in total duration of EEG seizures on day 14 Adverse events (any; serious; thought to be related to treatment; leading to discontinuation) | |

³ EEG: electroencephalogram; LEV: levetiracetam; RCT: randomised controlled trial

4 Table 5: Summary of included studies. Comparison 4: lamotrigine versus placebo

| Study | Population | Intervention | Comparison | Outcomes |
|------------|-------------|--------------|----------------|--------------------------------------|
| Frank 1999 | N=45 people | Lamotrigine | <u>Placebo</u> | Remained seizure |

2

| with absence seizures recruited; n=29 randomised after dose escalation and took part in dose escalation phase. n=28 people with absence seizures analysed (1 patient in lamotrigine group withdrew consent). Mean age: LTG: 6.9 years (SD 2.3) N=14 n=14 n=14 Patients who became seizure free after dose escalation phase were randomised to LTG or placebo LTG or placebo LTG or placebo LTG weeks. Placebo lasted 4 weeks free at end of placebo controlled phase Patients who became seizure free after dose escalation phase were randomised to LTG or placebo LTG or placebo LTG was tapered over 2 weeks. Placebo lasted 4 weeks free at end of placebo controlled phase Patients who became seizure free after dose escalation phase were randomised to LTG or placebo LTG was tapered over 2 weeks. Placebo lasted 4 weeks free at end of placebo controlled phase Adverse events - leading to withdrawal from study from study From the free at end of placebo controlled phase Adverse events - leading to withdrawal from study From the free at end of placebo controlled phase Adverse events - leading to withdrawal from study From the free at end of placebo controlled phase Adverse events - leading to withdrawal from study | Study | Population | Intervention | Comparison | Outcomes |
|---|-------|---|---|---|---|
| years (SD 3.1) FEG: electroencephalogram: LTG: lamotrigine: RCT: randomised controlled trial: SD: standard deviation | US | seizures recruited; n=29 randomised after dose escalation and took part in dose escalation phase. n=28 people with absence seizures analysed (1 patient in lamotrigine group withdrew consent). Mean age: LTG: 6.9 years (SD 2.3) Placebo: 8.8 years (SD 3.1) | Initial dose of 0.5mg/kg/day for 2 weeks, followed by 1mg/kg/day for 2 weeks. Following increases by 1mg/kg/ day according to response, maximum dose of 15mg/kg/day. Chewable 5mg, 25 and 100mg tables | Patients who became seizure free after dose escalation phase were randomised to LTG or placebo – LTG was tapered over 2 weeks. Placebo lasted 4 weeks | cebo controlled phase • Adverse events - leading to with-drawal from study |

EEG: electroencephalogram; LTG: lamotrigine; RCT: randomised controlled trial; SD: standard deviation

Table 6: Summary of included studies. Comparison 5: ethosuximide versus lamotrigine

| Study | Population | Intervention | Comparison | Outcomes |
|--|--|--|---|--|
| Glauser 2010 Multi-centre RCT US Reports 20 week follow-up data | N = 453 people with absence sei- zures (three- arm trial) Mean age: not provided | Ethosuximide n=156 250mg capsules or 250 mg per 5 ml of syrup. Highest allowable daily dose of 60mg/kg (to a maximum of 2000mg). | Lamotrigine n=149 5mg and 25mg chewable tablets or 25mg tablets. Highest allowable daily dose of 12mg/kg (to a maximum of 600 mg. | Freedom from treatment failure Adverse events (serious adverse events; intolerable adverse effects) Study drug discontinued Withdrawal from study Attentional dysfunction (proxy outcome for neuropsychological changes) |
| Glauser 2013 See Glauser 2010 Reports 12 month follow-up data | See Glauser 2010 | See Glauser 2010 | See Glauser 2010 | Freedom from treatment failure Adverse events (serious adverse events; intolerable adverse effects) Attentional dysfunction (proxy outcome for neuropsychological changes) |

| Study | Population | Intervention | Comparison | Outcomes |
|-------|------------|--------------|------------|---|
| | | | | Withdrawal from study |

- 1 RCT: randomised controlled trial
- 2 See the full evidence tables in appendix D and forest plots in appendix E.

3 Summary of the evidence

- 4 Across all the comparisons identified in this review, the majority identified a clinically im-
- 5 portant difference in at least one of the outcomes reported. For example, the comparison
- 6 lamotrigine versus sodium valproate showed a clinically important benefit in relation to the
- 7 number of people who were seizure free at 1 month and at 3 months and received sodium
- 8 valproate, and in relation to scores on the Conners' Continuous Performance Test, which fa-
- 9 voured those who received lamotrigine. When compared to placebo, those who received le-
- 10 vetiracetam showed an improved EEG response on days 11 to 14. When compared to pla-
- 11 cebo, lamotrigine showed a clinically important benefit in relation to the number of patients
- who were seizure free at 4 weeks. The comparison ethosuxumide versus lamotrigine showed
- a clinically important benefit in relation to the number of patients who were seizure free at 16
- or 20 weeks or at 12 months for those who received ethosuxumide.
- 15 There is a high level of uncertainty in the data presented and these findings should not there-
- 16 fore be considered to be definitive results. This uncertainty comes from the quality of the evi-
- dence, with the majority of results being assessed as very low, low or moderate quality. Out-
- 18 comes were most often downgraded due to risk of bias, with limited information provided re-
- 19 garding randomisation and allocation concealment. Outcomes were also downgraded due to
- 20 imprecision and the majority of studies only included a small number of participants; further
- 21 limiting confidence in the data.
- 22 A small number of outcomes were rated as being of high quality, indicating that the true ef-
- 23 fect for these treatments are similar to those calculated in this review. This was the case for
- the comparison of ethosuximide versus sodium valproate, for which there was high quality
- 25 evidence that sodium valproate was more effective than ethosuximide in relation to im-
- 26 provements in the symptoms of attention deficit disorder (Conners' Continuous Performance
- 27 Test score > 0.60) at 12 month follow-up. There was also high quality evidence showing an
- 28 important difference for sodium valproate compared to lamotrigine in relation to improve-
- ments in the symptoms of attention deficit disorder at the 16 or 20 week follow-up point; as
- well as freedom from treatment failure at the 16 or 20 week follow-up and the 12 month fol-
- 31 low-up. There was also high quality evidence indicating that ethosuximide provides important
- 32 benefits (when compared to lamotrigine) in relation to freedom from treatment failure at 16 or
- 33 20 weeks and at 12 months.

34 35

- Uncertainty in the data is further exacerbated by the absence of evidence on health-related
- quality of life, as well as a number of antiseizure therapies included in the review protocol.

37 Quality assessment of clinical outcomes included in the evidence review

- 38 See the clinical evidence profiles in appendix F.
- 39 Economic evidence
- 40 Included studies
- 41 A single economic search was undertaken for all topics included in the scope of this guide-
- 42 line. See supplementary material 2 for details.

1 Excluded studies

- 2 A single economic search was undertaken for all topics included in the scope of this guide-
- 3 line. See supplementary material 2 for details.

4 Summary of studies included in the economic evidence review

5 No studies were identified which were applicable to this review question.

6 Economic model

- 7 No economic modelling was undertaken for this review because the committee agreed that
- other topics were higher priorities for economic evaluation. 8

9 The committee's discussion of the evidence

10 Interpreting the evidence

11 The outcomes that matter most

- 12 The committee agreed that seizure freedom should be a critical outcome for this review as
- 13 this is considered to be the main objective of treatment for children with absence seizures.
- 14 24-hour EEG seizure freedom was also included as a critical outcome as absence seizures
- show a specific pattern on EEG which usually improves or resolves with effective treatments. 15
- 16 The committee also agreed that time to discontinuation of treatment or change of medication,
- and adverse events should be included as critical outcomes to ensure that data on treatment 17
- 18 acceptability and tolerability were included.
- 19 Neuropsychological changes and health related quality of life were also included as im-
- 20 portant outcomes as deterioration in these may suggest adverse treatment effects.

21 The quality of the evidence

- 22 The committee were presented with data on 5 different comparisons, however, meta-analysis
- 23 was only possible for 1 outcome in 1 comparison due to variation in the interventions and
- 24 measurement of outcomes. All data presented related to children and young people under
- the age of 18 years. The quality of the evidence for this review was assessed using GRADE 25
- methodology. The majority of outcomes were considered very low, low or moderate quality 26
- 27 indicating uncertainty in the reliability of the data. This was with the exception of some of the
- outcomes reported by Glauser 2010 and Glauser 2013, which were considered high quality; 28
- 29 mainly as a result of a low risk of bias and more precise estimates. Outcomes were most of-
- ten downgraded due to risk of bias, with limited information provided regarding randomisation 30
- and allocation concealment. Outcomes were also downgraded due to imprecision and the 31 32 majority of studies only included a small number of participants; further limiting confidence in
- the data. 33
- 34 No studies reported on the outcome health-related quality of life.

35 Benefits and harms

- The committee used the evidence presented and their clinical knowledge and expertise to 36
- 37 make recommendations.
- 38 No evidence was identified which reported on the effectiveness of ASMs as treatment for ab-
- sence seizures in people over 18 years; however, the committee agreed that, for recommen-39
- dations on adults, it was appropriate to extrapolate from the evidence on children and young 40
- 41 people given the similar pathophysiology in both age groups.

- 1 The committee agreed that, prior to starting antiseizure therapy there should be a discussion
- with the person, their family and carers, if appropriate, about an strategy according to their
- 3 seizure type, treatment goals and the preferences of the person and their family or carers, as
- 4 appropriate. Treatment plans should be regularly reassessed, and its agreement should in-
- 5 clude a transparent explanation of the seizure type, severity and duration of adverse effects
- 6 that the person with epilepsy may experience and how should these be managed. The per-
- son, their family and carers, should also be made aware that they should be taking the least
- 8 amount of medicines as possible to be effective due to the side effects of being on numerous
- 9 medications.

48

- 10 The committee emphasised that monotherapy should be used in the first instance. When
- starting alternative antiseizure medications, the dose of the new antiseizure medication
- should be slowly increased, whilst the existing antiseizure medication is tapered off. When
- 13 starting add-on antiseizure medications, the additional antiseizure medication should be
- carefully titrated, in line with the BNF guidance, adverse events monitored, and there should
- 15 be a frequent treatment review.

Absence seizures (including childhood absence epilepsy)

- 17 There was some evidence which indicated that ethosuximide improves freedom from treat-
- ment failure in people with absence seizures (including childhood absence epilepsy). Alt-
- 19 hough this evidence was generally of low quality, there was some high quality evidence
- 20 which also showed that ethosuximide had an important benefit in relation to this outcome
- 21 (when compared to lamotrigine, at 16/20 weeks follow-up, and at 12 months follow-up). The
- 22 committee therefore agreed that this should be a strong recommendation, particularly as
- 23 ethosuximide is the most widely used ASM in the treatment of absence seizures (including
- childhood absence epilepsy) and is associated with seizure remission.
- 25 Evidence was also identified which suggested that sodium valproate was effective in the
- 26 treatment of absence seizures (including childhood absence epilepsy). Although the evi-
- 27 dence was generally of low quality, the committee agreed to make a recommendation stating
- that sodium valproate can be offered as second-line alternative or add-on treatment given its
- association with extended seizure freedom (when compared to lamotrigine). Furthermore,
- 30 the committee highlighted that sodium valproate is commonly used in clinical practice. The
- 31 committee discussed in detail the risks associated with use of sodium valproate in women
- 32 and girls who are able to have children. They agreed to include in the recommendation the
- caveat that sodium valproate should only be offered to girls if they are under the age of 10,
- 34 noting that girls over this age who are approaching puberty and still experiencing absence
- 35 seizures are likely to have their diagnosis and treatment reviewed.
- There was some evidence which suggested that lamotrigine and levetiracetam were effective
- in the treatment of absence seizures (including childhood absence epilepsy). The committee
- 38 decided to recommend these as third-line treatments because ethosuxumide showed to have
- 39 better outcomes than lamotrigine, including better seizure control. Compared to sodium
- valproate, lamotrigine showed to be less effective for seizure control, although there were not
- 41 differences between both ASMs for adverse events and treatment withdrawal. Furthermore,
- 42 when lamotrigine was compared with placebo, it showed to be more effective for seizure con-
- 43 trol and no adverse events were reported in neither of the trial arms. There was a placebo-
- 44 controlled trial for levetiracetam which showed better seizure control in people who received
- levetiracetam and no clinically important differences in terms of adverse events.
- The committee noted that although other antiseizure medications are used in clinical practice
- and may benefit some people, it should be highlighted that some can exacerbate seizures.

Absence seizures with other seizure types

- 49 For boys and men; girls aged under 10 years who are unlikely to need treatment when they
- are old enough to have children, and women who are unable to have children with absence

- 1 seizures with other seizure types, the committee agreed to recommend sodium valproate as
- 2 first-line treatment given its association with higher remission rates and tolerability in children
- 3 and young people.
- 4 As the evidence also indicated that lamotrigine and levetiracetam are effective, the commit-
- 5 tee agreed to recommend the use of these as second-line treatment if first-line is unsuccess-
- ful in boys and men with absence seizures and other seizure types, and as first-line treat-
- 7 ment in women and girls of childbearing age (given the risks associated with sodium
- 8 valproate). Lamotrigine and levetiracetam were recommended as second-line treatments be-
- 9 cause, compared to sodium valproate, lamotrigine showed to be less effective for seizure
- 10 control, although there were no differences between both ASMs for adverse events and
- 11 treatment withdrawal. Furthermore, when lamotrigine was compared with placebo, it showed
- to be more effective for seizure control and no adverse events were reported in neither of the
- trial arms. There was a placebo-controlled trial for levetiracetam, which showed better sei-
- zure control in people who received levetiracetam and no clinically important differences in
- 15 terms of adverse events
- 16 Based on the evidence identified, lamotrigine and levetiracetam were also recommended as
- an add-on treatment in boys and men with absence seizures and other seizure types.
- As there was evidence for ethosuxumide in children and young people, the committee
- agreed to include this drug as an add-on treatment if first-line treatment is unsuccessful. The
- 20 committee believed that ethosuxumide may be needed in some situations, for example, in
- 21 cases of absence seizures co-existing with other seizure types.
- 22 As there was evidence of effectiveness, the committee felt that it was necessary to make a
- 23 recommendation stating that sodium valproate may be prescribed to women and girls of
- 24 childbearing age, however they agreed that the recommendation should emphasise that this
- should be seen as a 'last resort' and must be done only after discussion of the risks and
- benefits. If sodium valproate is prescribed to women and girls able to have children, clini-
- 27 cians must follow MHRA guidance, which includes ensuring the continuous use of highly ef-
- 28 fective contraception and the enrolment of the girl or woman in a pregnancy prevention pro-
- 29 gramme, if appropriate.
- The committee noted that although other antiseizure medications are used in clinical practice
- and may benefit some people, it should be highlighted that some can exacerbate seizures.

32 Cost effectiveness and resource use

- 33 A systematic review of the economic literature was conducted but no relevant studies were
- identified which were applicable to this review question.
- 35 These recommendations represent current practice and therefore no resource impact is an-
- 36 ticipated. Other than levetiracetam where a third line treatment is needed, the recommended
- 37 antiseizure medications are identical to those of the previous NICE guideline. It is not antici-
- 38 pated that there will be any change in the use of levetiracetam as a result of these recom-
- mendations as it is already used in clinical practice in such circumstances.

40 Other factors the committee took into account

- In line with the MHRA, the committee emphasised that long-term treatment with sodium
- 42 valproate can cause decreased bone mineral density and increased risk of osteomalacia.
- The committee noted that appropriate supplementation should be considered for those at
- 44 risk.

45 Recommendations supported by this evidence review

- This evidence review supports recommendations 5.3.1-5.3.9.
- 47 15

References

2

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- 18 Casale, E. J., Lamictal (lamotrigine) monotherapy for typical absence seizures in children,
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24 Glauser 2010

- 25 Glauser, T. A., Cnaan, A., Shinnar, S., Hirtz, D. G., Dlugos, D., Masur, D., Clark, P. O., Cap-
- parelli, E. V., Adamson, P. C., Ethosuximide, valproic acid, and lamotrigine in childhood ab-
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28 **Sato 1982**

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- 30 sence seizures, Neurology, 32, 157-163, 1982

Appendices

2 Appendix A – Review protocols

- 3 Review protocol for review question: What antiseizure therapies (monotherapy or add-on) are effective in the treatment of
- 4 absence seizures?

5 Table 7: Review protocol

| Field | Content |
|------------------------------|--|
| PROSPERO registration number | CRD42019159420 |
| Review title | Effectiveness of antiseizure therapies for absence seizures |
| Review question | What antiseizure therapies (monotherapy or add-on) are effective in the treatment of absence seizures? |
| Objective | The objective of this review is to determine which antiseizure therapies improve outcomes in people with epilepsy who have absence seizures. This review will determine the effectiveness of antiseizure therapies given alone (monotherapy) or as add-ons (combination therapy). |
| Searches | The following databases will be searched: CDSR CENTRAL DARE HTA MEDLINE & MEDLINE In-Process and Other Non-Indexed Citations Embase EMCare Searches will be restricted by: |

| Field | Content |
|---|--|
| | Date: No date limit |
| | English language sutdies |
| | Human studies |
| | RCT and systematic review study design filter |
| Condition or domain being studied | Epilepsy with absence seizures |
| Population | Inclusion |
| | People with confirmed epilepsy with absence seizures |
| | Exclusion |
| | Newborn babies (under 28 days) with acute symptomatic seizures |
| Intervention/Exposure/Test | The following antiseizure therapies and their combinations will be considered: |
| | Acetazolamide |
| | • Clobazam |
| | Clonazepam |
| | Ethosuximide |
| | Ketogenic diet (included as this is an accepted first or second line treatment for these type of seizures) |
| | Lamotrigine |
| | Levetiracetam |
| | Methsuximide/mesuximide |
| | Sodium Valproate |
| | Topiramate |
| | Zonisamide |
| Comparator/Reference standard/Confounding | Any of the above and their combinations |
| factors | No treatment/placebo |
| Types of study to be included | Systematic review of RCTs |
| | • RCTs |

| Field | Content |
|---|--|
| Other exclusion criteria | Studies with a mixed population (this is, including children and young people with epilepsy and others with a condition different to epilepsy) will be excluded, unless subgroup analysis for epilepsy has been reported Studies with a mixed population (this is, including people with epilepsy with different seizure types) will be excluded, unless subgroup analysis for epilepsy with absence seizures has been reported. Conference abstracts will be excluded because these do not typically provide sufficient information to fully assess risk of bias |
| Context | Recommendations will apply to those receiving care in any healthcare settings (for example, community, primary, secondary care) |
| Primary outcomes (critical outcomes) | Seizure freedom (12 months data and short term, (minimum 3 months with 100% freedom) of starting treatment) Due to anticipated heterogeneity in reporting of seizure freedom, data will be extracted as presented within included studies. Where a study reports multiple variants then all data will be extracted. For decision making priority will be given to data presented as "time to 12 months seizure freedom", (this is, time to event: HR or mean time) followed by "achievement of 12 months seizure freedom" (RR). Minimum follow up data of 3 months will be included. 24 hour EEG seizure freedom Time to withdrawal of treatment or change of medication (for example, because of uncontrollable seizures) Adverse events, as assessed by: % of patients with reported side effects (trial defined adverse and serious adverse effects) treatment cessation due to adverse event (dichotomous outcome only) mortality |
| Secondary outcomes (important outcomes) | Neuropsychological changes (IQ testing, or other validated tools) |

| Field | Content |
|--|--|
| | Health-related overall quality of life (measured using validated tools only) Outcomes are in line with those described in the core outcome set for epilepsy http://www.cometinitiative.org/studies/searchresults |
| Data extraction (selection and coding) | All references identified by the searches and from other sources will be uploaded into STAR and de-duplicated. |
| | The full text of potentially eligible studies will be retrieved and will be assessed in line with the inclusion criteria. Duplicate screening will not be undertaken for this review question. A standardised form will be used to extract data from studies (see Developing NICE guidelines: the manual section 6.4) and will include: study setting; design; aim; study dates; funding; sample size; participant demographics and baseline characteristics; inclusion and exclusion criteria; details of intervention and controls; study methodology; recruitment and study completion rates; outcomes and times of measurement; and information for assessment of risk of bias. Full versions of the selected studies will be obtained for assessment. Studies that fail to meet the inclusion criteria, once the full version has been checked, will be excluded at this stage. Each study excluded after checking the full version will be listed, along with the reasons for its exclu- |
| | sion. All data extraction will be quality assured by a senior reviewer. Draft included and excluded studies tables will be circulated to the Topic Group for their comments. Resolution of disputes will be by discussion between the senior reviewer, Topic Advisor and Chair. |
| Risk of bias (quality) assessment | Quality assessment of individual studies will be performed using the following checklists: ROBIS tool for systematic reviews Cochrane RoB tool v.2 for RCTs The quality assessment will be performed by one reviewer and this will be quality assessed by a senior reviewer. |
| Strategy for data synthesis | Depending on the availability of the evidence, the findings will be summarised narratively or quantitatively. <u>Data synthesis</u> |

| Field | Content |
|-------|---|
| | Where possible pairwise meta-analyses will be conducted using Cochrane Review Manager software. A fixed effect meta-analysis will be conducted and data will be presented as risk ratios for dichotomous outcomes. Peto odds ratio will be used for outcomes with zero events in one arm and <1% events in the other. Risk difference will be used for outcomes with zero events in both arms. Mean differences or standardised mean differences will be presented for continuous outcomes. |
| | Heterogeneity Heterogeneity in the effect estimates of the individual studies will be assessed using the I2 statistic. I² values of greater than 50% and 75% will be considered as significant and very significant heterogeneity, respectively. |
| | In the presence of heterogeneity, sub-group analysis will be conducted: according to the risk of bias of individual studies study location |
| | Exact sub-group analysis may vary depending on differences identified within included studies |
| | If heterogeneity cannot be explained using these methods, random effects model will be used. If heterogeneity remains above 75% and cannot be explained by sub-group analysis. Reviewers will consider if meta-analysis is appropriate given characteristics of included studies. |
| | Minimal important differences (MIDs): |
| | Default MIDs will be used for risk ratios and continuous outcomes only, unless the committee prespecifies published or other MIDs for specific outcomes |
| | • For risk ratios: 0.8 and 1.25. |
| | For continuous outcomes: |
| | o For one study: the MID is calculated as +/-0.5 times the baseline SD of the control arm. |
| | For two studies: the MID is calculated as +/-0.5 times the mean of the SDs of the control arms at baseline. If baseline SD is not available, then SD at follow up will be used. |
| | o For three or more studies (meta-analysed): the MID is calculated by ranking the studies in or- |

| Field | Content | | |
|---|---|--|--|
| | der of SD in the control arms. The MID is calculated as +/- 0.5 times median SD. o For studies that have been pooled using SMD (meta-analysed): +0.5 and -0.5 in the SMD scale are used as MID boundaries. Validity The confidence in the findings across all available evidence will be evaluated for each outcome. | | |
| | using an adaptation of the 'Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international GRADE working group: http://www.gradeworkinggroup.org/ | | |
| Analysis of sub-groups (stratification) | Stratification | | |
| | | sults will be presented separately by: | |
| | Those with and without learning difficulties/disabilities Age of onset (≤10 years and > 10 years old) | | |
| | Eyelid myoclonia | | |
| | Photosensitivity | | |
| | Typical and atypical absence seizures | | |
| Type and method of review | \boxtimes | Intervention | |
| | | Diagnostic | |
| | | Prognostic | |
| | | Qualitative | |
| | | Epidemiologic | |
| | | Service Delivery | |
| | | Other (please specify) | |
| Language | English | | |
| Country | England | | |
| Anticipated or actual start date | 20 February 2020 | | |
| Anticipated completion date | 21 April 2021 | | |

| Field | Content | | |
|--|---|----------|-----------|
| Stage of review at time of this submission | Review stage | Started | Completed |
| | Preliminary searches | V | • |
| | Piloting of the study selection process | V | V |
| | Formal screening of search results against eligibility criteria | V | |
| | Data extraction | V | ~ |
| | Risk of bias (quality) assessment | V | V |
| | Data analysis | V | V |
| Named contact | 5a. Named contact National Guideline Alliance 5b Named contact e-mail epilepsies@nice.org.uk 5e Organisational affiliation of the review National Institute for Health and Care Excellence (NICE) and National Guideline Alliance | | |
| Review team members | The National Guideline Alliance technical team | | |
| Funding sources/sponsor | This systematic review is being completed by the National Guideline Alliance which receives funding from NICE. | | |
| Conflicts of interest | All guideline committee members and anyone who has direct input into NICE guidelines (including the evidence review team and expert witnesses) must declare any potential conflicts of interest in line with NICE's code of practice for declaring and dealing with conflicts of interest. Any relevant interests, or changes to interests, will also be declared publicly at the start of each guideline committee meeting. Before each meeting, any potential conflicts of interest will be considered by the guideline committee Chair and a senior member of the development team. Any decisions to exclude a person from all or part of a meeting will be documented. Any changes to a member's declaration of interests will be recorded in the minutes of the meeting. Declarations of interests will be published with the final guideline. | | |

| Field | Content | |
|--|---|--|
| Collaborators | Development of this systematic review will be overseen by an advisory committee who will use the review to inform the development of evidence-based recommendations in line with section 3 of Developing NICE guidelines: the manual . Members of the guideline committee are available on the NICE website: https://www.nice.org.uk/guidance/indevelopment/gid-ng10112 | |
| Other registration details | Not applicable | |
| URL for published protocol | https://www.crd.york | .ac.uk/prospero/display_record.php?RecordID=159420 |
| Dissemination plans | NICE may use a range of different methods to raise awareness of the guideline. These include standard approaches such as: notifying registered stakeholders of publication publicising the guideline through NICE's newsletter and alerts issuing a press release or briefing as appropriate, posting news articles on the NICE website, using social media channels, and publicising the guideline within NICE. | |
| Keywords | Epilepsy; Absence seizures | |
| Details of existing review of same topic by same authors | Not applicable | |
| Current review status | \boxtimes | Ongoing |
| | | Completed but not published |
| | | Completed and published |
| | | Completed, published and being updated |
| | | Discontinued |
| Additional information | Not applicable | |
| Details of final publication | www.nice.org.uk | |

1 Appendix B - Literature search strategies

2 Literature search strategies for review question: What antiseizure therapies3 (monotherapy or add-on) are effective in the treatment of absence seizures?

Clinical

6

4 5

7

8 9

10

Database(s): EMCare, MEDLINE and Embase (Multifile) – OVID

EMCare 1995 to December 03, 2019; Embase Classic+Embase 1947 to 2019 December 03; Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily 2019 December 03, 2019

Date of last search: 03 December 2019

11 12 13

Multifile database codes: emcr=EMCare; emczd=Embase Classic+Embase; ppez= MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily

| # | searches |
|----|--|
| 1 | (seizure and absence).sh. use emczd, emcr or seizures/ use ppez or ((absence adj2 (convulsion* or seizure*)) or ((typical or atypical) adj absenc*) or petit mal* or pyknolepsy or typical absence*).ti,ab. |
| 2 | ethosuximide/ use emczd, emcr or ethosuximide/ use ppez or (emeside or ethosuccimid* or ethosuccinimid* or ethosuximide or ethylmethylsuccimide or ethylsuximide or ethymal or etosuximida or mesentol or pemal or petimid or petinimid* or petnidan or pyknolepsin or pyknolepsinum or ronton or simatin or succinutin or sucsilep or suksilep or suxilep or suximal or suxinutin or zarondan or zarontin).ti,ab. |
| 3 | fat intake/ or glycemic index/ or ketogenic diet/ or exp low carbohydrate diet/ or exp triacylglycerol/ |
| 4 | 3 use emczd, emcr |
| 5 | exp diet, carbohydrate-restricted/ or exp dietary fats/ or glycemic index/ or exp triglycerides/ |
| 6 | 5 use ppez |
| 7 | ((adequate adj3 protein*) or atkin* or keto* or kd* or (carbohydrate* adj5 (restrict* or low* or reduc*)) or ((glycemic or glycaemic) adj5 (index or treat* or modulat*)) or (high fat* adj5 (diet* or plan* or treat*)) or keto or ketogenic or ketogenous or ketotic or low carb* or lchf or low glyc* index treatment* or lgit or (medium chain adj (tryglyceride* or triglyceride*)) or mct*).ti,ab. |
| 8 | or/4,6-7 |
| 9 | lamotrigine/ use emczd, emcr or lamotrigine/ use ppez or (crisomet or labileno or lamepil or lamictal or lamictin or lamotkal or lamodex or lamogine or lamotrigin* or lamotrix or neurium).ti,ab. |
| 10 | levetiracetam/ use emczd, emcr, ppez or (elepsia or keppra or kopodex or levetiracetam* or matever or spritam).ti,ab. |
| 11 | topiramate/ use emczd, emcr, ppez or (epitomax or topamax or topiramate or acomicil or ecuram or epiramat or epitomax or epitoram or erravia or etopro or fagodol or jadix or lusitrax or maritop or oritop or piraleps or pirantal or pirepil or qudexy or ramas or sincronil or talopam or tiramat or topaben or topamac or topamax or topepsil or topibrain or topilek or topimark or topimax or topiramat* or topiramat or topiratore or topit or toramat or torlepta or trokendi).ti,ab. |
| 12 | valproic acid/ use emczd, emcr, ppez or (convulsofin or delepsine or depacon* or depaken* or depakin* or depakente or depa |
| 13 | zonisamide/ use emczd, emcr or zonisamide/ use ppez or (excegran or excemid or zonegran or zonisamid*).ti,ab. |
| 14 | clobazam/ use emczd, emcr or clobazam/ use ppez or (chlorepin or chlorepine or clobazam or clobazem or clorepin or frisium or noiafren or onfi or urbadan or urbanil or urbanyl).ti,ab. |
| 15 | clonazepam/ use emczd, emcr or clonazepam/ use ppez or (aklonil or antelepsin or clonazepam or clonex or clonopam or clonopin or clonotril or coquan or iktorivil or kenoket or klonazepam or klonopin |

| # | searches |
|----------|--|
| | or kriadex or landsen or lonazep or paxam or povanil or ravotril or rivatril or rivotril).ti,ab. |
| 16 | mesuximide/ use emczd, emcr or (alpha methylphensuximide or celontin or methosuximide or celontine or mesuximide or methylsuximide or metsuccimide or petinutin).ti,ab. |
| 17 | acetazolamide/ use emczd, emcr or acetazolamide/ use ppez or (acetadiazol or acetamox or acetazol amide or acetazolam or acetazolamid* or acetazolamine or acetazolamid* or acetazolamine or ak zol or akzol or albox or apoacetazolamide or azetazolamide or carbinib or carbonic anhydrase inhibitor or cidamex or dazamide or defiltran or dehydratin or diacarb or diamox or diluran or diomax or diuramid* or diutazol or edemox or eumicton or fonurit or genephamide or glaucomed* or glauconox or glaupax or huma zolamide or humazolamide or ledamox or lediamox or ledimox or natrionex or nephramid or novozolamide or storzolamide or ulcosilvanil or ulcosylvanil).ti,ab. |
| 18 | or/2,8-17 |
| 19 | clinical trials as topic.sh. or (controlled clinical trial or pragmatic clinical trial or randomized controlled trial).pt. or (placebo or randomi#ed or randomly).ab. or trial.ti. |
| 20 | 19 use ppez |
| 21 | (controlled clinical trial or pragmatic clinical trial or randomized controlled trial).pt. or drug therapy.fs. or (groups or placebo or randomi#ed or randomly or trial).ab. |
| 22 | 21 use ppez |
| 23 | crossover procedure/ or double blind procedure/ or randomized controlled trial/ or single blind procedure/ or (assign* or allocat* or crossover* or cross over* or ((doubl* or singl*) adj blind*) or factorial* or placebo* or random* or volunteer*).ti,ab. |
| 24 | 23 use emczd, emcr |
| 25 | or/20,22,24 |
| 26 | meta-analysis/ |
| 27 | meta-analysis as topic/ or systematic reviews as topic/ "systematic review"/ |
| 28 29 | meta-analysis/ |
| 30 | (meta analy* or metanaly* or metaanaly*).ti,ab. |
| 31 | ((systematic or evidence) adj2 (review* or overview*)).ti,ab. |
| 32 | ((systematic* or evidence*) adj2 (review* or overview*)).ti,ab. |
| 33 | (reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab. |
| 34 | (search strategy or search criteria or systematic search or study selection or data extraction).ab. |
| 35 | (search* adj4 literature).ab. |
| 36 | (Medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab. |
| 37 | cochrane.jw. |
| 38 | ((pool* or combined) adj2 (data or trials or studies or results)).ab. |
| 39 | (or/26-27,30,32-38) use ppez |
| 40 | (or/28-31,33-38) use emczd, emcr |
| 41 42 | or/39-40 or/25,41 |
| 42 | 1 and 18 and 42 |
| 44 | limit 43 to english language |
| 45 | ((letter.pt. or letter/ or note.pt. or editorial.pt. or case report/ or case study/ or (letter or comment*).ti.) not (randomized controlled trial/ or random*.ti,ab.)) or ((animal/ not human/) or nonhuman/ or exp animal experiment/ or exp experimental animal/ or animal model/ or exp rodent/ or (rat or rats or mouse or mice).ti.) |
| 46 | 45 use emez |
| 47 | ((letter/ or editorial/ or news/ or exp historical article/ or anecdotes as topic/ or comment/ or case report/ or (letter or comment*).ti.) not (randomized controlled trial/ or random*.ti,ab.)) or ((animals not humans).sh. or exp animals, laboratory/ or exp animal experimentation/ or exp models, animal/ or exp rodentia/ or (rat or rats or mouse or mice).ti.) |
| 48 | 47 use mesz |
| 49 | 46 or 48 |
| 50 | 44 not 49 |
| | |

4

Database(s): Cochrane LibraryCochrane Database of Systematic Reviews, Issue 12 of 12, December 2019; Cochrane Central Register of Controlled Trials, Issue 12 of 12, December 2019

1 2 Date of last search 03 December 2019

| ш | anavahan | | | | | |
|----------------|---|--|--|--|--|--|
| # #1 | searches mesh descriptor: [seizures] explode all trees | | | | | |
| #1 | (((absence near/2 (convulsion* or seizure*)) or ((typical or atypical) near/1 absenc*) or "petit mal*" or | | | | | |
| π ∠ | pyknolepsy or "typical absence*")):ti,ab,kw | | | | | |
| #3 | #1 or #2 | | | | | |
| #4 | mesh descriptor: [ethosuximide] this term only | | | | | |
| #5 | ((emeside or ethosuccimid* or ethosuccinimid* or ethosuximide or ethylmethylsuccimide or | | | | | |
| | ethylsuximide or ethymal or etosuximida or mesentol or pemal or petimid or petinimid* or petnidan or | | | | | |
| | pyknolepsin or pyknolepsinum or ronton or simatin or succinutin or sucsilep or suksilep or suxilep or | | | | | |
| | suximal or suxinutin or zarondan or zarontin)):ti,ab,kw | | | | | |
| #6 | mesh descriptor: [diet, carbohydrate-restricted] explode all trees | | | | | |
| #7 | mesh descriptor: [dietary fats] explode all trees | | | | | |
| #8 | mesh descriptor: [glycemic index] this term only | | | | | |
| #9 | mesh descriptor: [triglycerides] this term only | | | | | |
| #10 | (((adequate near/3 protein*) or atkin* or keto* or kd* or (carbohydrate* near/5 (restrict* or low* or reduc*)) or ((glycemic or glycaemic) near/5 (index or treat* or modulat*)) or ("high fat*" near/5 (diet* or | | | | | |
| | plan* or treat*)) or keto or ketogenic or ketogenous or ketotic or "low carb*" or lchf or "low glyc* index | | | | | |
| | treatment*" or lgit or ("medium chain" near/1 (tryglyceride* or triglyceride*)) or mct*)):ti,ab,kw | | | | | |
| #11 | mesh descriptor: [lamotrigine] this term only | | | | | |
| #12 | ((crisomet or labileno or lamepil or lamictal or lamictin or lamiktal or lamodex or lamogine or lamotrigin* | | | | | |
| | or lamotrix or neurium)):ti,ab,kw | | | | | |
| #13 | mesh descriptor: [levetiracetam] this term only | | | | | |
| #14 | ((elepsia or keppra or kopodex or levetiracetam* or matever or spritam)):ti,ab,kw | | | | | |
| #15 | mesh descriptor: [topiramate] this term only | | | | | |
| #16 | ((epitomax or topamax or topiramate or acomicil or ecuram or epiramat or epitomax or epitoram or er- | | | | | |
| | ravia or etopro or fagodol or jadix or lusitrax or maritop or oritop or piraleps or pirantal or pirepil or | | | | | |
| | qudexy or ramas or sincronil or talopam or tiramat or topaben or topamac or topamax or topepsil or topibrain or topilek or topimark or topimax or topiramat* or topiramato or topiratore or topit or toramat or | | | | | |
| | torlepta or trokendi)):ti,ab,kw | | | | | |
| #17 | mesh descriptor: [valproic acid] this term only | | | | | |
| #18 | ((convulsofin or delepsine or depacon* or depaken* or depakin* or depakote or depalept or deprakine | | | | | |
| | or "di n propylacetate" or "di n propylacetate sodium" or "di n propylacetic acid" or diplexil or "dipropyl | | | | | |
| | acetate" or "dipropyl acetic acid" or dipropylacetate or "dipropylacetate sodium" or "dipropylacetatic | | | | | |
| | acid" or "dipropylacetic acid" or diprosin or divalproex or epilam or epilex or "epilim chrono" or "epilim | | | | | |
| | chromosphere" or "epilim enteric" or epilim or episenta or "epival cr" or ergenyl or "espa valept" or eve- | | | | | |
| | riden or goilim or hexaquin or labazene or leptilan or leptilanil or micropakine or mylproin or "myproic acid" or "n dipropylacetic acid" or orfiril or orlept or petilin or "propylisopropylacetic acid" or | | | | | |
| | propymal or "semisodium valproate" or "sodium 2 propylpentanoate" or "sodium 2 propylvalerate" or | | | | | |
| | "sodium di n propyl acetate" or "sodium di n propylacetate" or "sodium dipropyl acetate" or "sodium di- | | | | | |
| | propylacetate" or "sodium n dipropylacetate" or stavzor or "valberg pr" or valcote or valepil or valeptol or | | | | | |
| | valerin or "valhel pr" or valoin or valpakine or valparin or valporal or valprax or valpro or valproate or | | | | | |
| | valprodura or "valproic acid" or valprosid or valprotek or valsup or vupral)):ti,ab,kw | | | | | |
| #19 | mesh descriptor: [zonisamide] this term only | | | | | |
| #20 | ((excegran or excemid or zonegran or zonisamid*)):ti,ab,kw | | | | | |
| #21 | mesh descriptor: [clobazam] this term only | | | | | |
| #22 | ((chlorepin or chlorepine or clobazam or clobazepam or clorepin or frisium or noiafren or onfi or urba- | | | | | |
| | dan or urbanil or urbanyl)):ti,ab,kw | | | | | |
| #23 | mesh descriptor: [clonazepam] this term only | | | | | |
| #24 | ((aklonil or antelepsin or clonazepam or clonex or clonopam or clonopin or clonotril or coquan or iktorivil | | | | | |
| | or kenoket or klonazepam or klonopin or kriadex or landsen or lonazep or paxam or povanil or ravotril | | | | | |
| "0 5 | or rivatril or rivotril)):ti,ab,kw | | | | | |
| #25 | (("alpha methylphensuximide" or celontin or methosuximide or celontine or mesuximide or methylphensuximide or | | | | | |
| #26 | methsuximide or methylsuximide or metsuccimide or petinutin)):ti,ab,kw mesh descriptor: [acetazolamide] this term only | | | | | |
| #27 | ((acetadiazol or acetamox or "acetazol amide" or acetazolam or acetazolamid* or acetazolamine or | | | | | |
| π ∠ I | acetazoleamid* or acetozolamine or "ak zol" or akzol or albox or apoacetazolamide or azetazolamide or | | | | | |
| | carbinib or "carbonic anhydrase inhibitor" or cidamex or dazamide or defiltran or dehydratin or diacarb | | | | | |
| | or diamox or diluran or diomax or diuramid* or diutazol or edemox or eumicton or fonurit or genepha- | | | | | |
| | mide or glaucomed* or glauconox or glaupax or "huma zolamide" or humazolamide or ledamox or ledi- | | | | | |
| | amox or ledimox or natrionex or nephramid or novozolamide or storzolamide or ulcosilvanil or ulcosyl- | | | | | |
| # 00 | vanil)):ti,ab,kw | | | | | |
| #28 | {or #4-#27} | | | | | |
| #29 | #3 and #28 | | | | | |

4

Database(s): Database of Abstracts of Reviews of Effects - CRD

Cochrane Database of Systematic Reviews, Issue 12 of 12, December 2019; Cochrane Central Register of Controlled Trials, Issue 12 of 12, December 2019

Date of last search 03 December 2019

5 6

| # | Searches |
|----|--|
| #1 | mesh descriptor seizures explode all trees |
| #2 | (((absence near2 (convulsion* or seizure*)) or ((typical or atypical) near1 absenc*) or "petit mal*" or pyknolepsy or "typical absence*")) |
| #3 | #1 or #2 |

7 8

Economic

9

11 12

Database(s): MEDLINE & Embase (Multifile) - OVID

Embase Classic+Embase 1947 to 2021 March 31; Ovid MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily 1946 to March 31, 2021 Date of last search: 31 March 2021

13 14

Multifile database codes: emczd=Embase Classic+Embase; ppez= MEDLINE(R) and Epub Ahead of Print, In-Process & Other Non-Indexed Citations and Daily

| # | searches | | | | | | |
|----|--|--|--|--|--|--|--|
| 1 | exp epilepsy/ or exp seizure/ or "seizure, epilepsy and convulsion"/ | | | | | | |
| 2 | 1 use emczd | | | | | | |
| 3 | exp epilepsy/ or seizures/ or seizures, febrile/ or exp status epilepticus/ | | | | | | |
| 4 | 3 use ppez | | | | | | |
| 5 | (epilep* or seizure* or convuls*).ti,ab. or (continous spike wave of slow sleep or infant* spasm*).ti,ab. | | | | | | |
| 6 | (seizure and absence).sh. use emczd, emcr or seizures/ use ppez or ((absence adj2 (convulsion* or seizure*)) or ((typical or atypical) adj absenc*) or petit mal* or pyknolepsy or typical absence*).ti,ab. | | | | | | |
| 7 | (atonic seizure or tonic seizure).sh. use emczd, emcr or exp seizures/ use ppez or ((drop or akinetic or atonic or tonic) adj2 (attack* or epileps* or seizure* or convulsion*)).ti,ab. or brief seizure.ti,ab. or (tonic adj3 atonic adj3 (attack* or epileps* or seizure* or convulsion*)).ti,ab. | | | | | | |
| 8 | exp benign childhood epilepsy/ use emczd, emcr or epilepsy, rolandic/ use ppez or (bcects or bects or brec or benign epilepsy or (benign adj2 (childhood or neonatal or pediatric or paediatric) adj2 epileps*) or (benign adj2 (childhood or neonatal or pediatric or paediatric) adj2 (convulsion* or epileps* or seizure* or spasm*)) or (benign adj3 (convulsion* or epileps*) adj2 centrotemporal adj2 spike*) or cects or ((centralopathic or centrotemporal or temporal-central focal) adj (convulsion* or epileps* or seizure*)) or ((osylvian or postrolandic or roland*) adj2 (convulsion* or epileps* or seizure* or spasm*))).ti,ab. | | | | | | |
| 9 | exp generalized epilepsy/ use emczd, emcr or exp epilepsy, generalized/ use ppez | | | | | | |
| 10 | (((akinetic or atonic or central or diffuse or general or generali?ed or idiopathic or tonic) adj3 (epilep* or seizure*)) or ((childhood absence or juvenile absence or myoclonic or myoclonia or myoclonic astatic or myoclonus or gtcs) adj2 epilep*) or (epilepsy adj2 eyelid myoclonia) or (ige adj2 phantom absenc*) or impulsive petit mal or (janz adj3 (epilep* or petit mal)) or jeavons syndrome* or ((janz or lafora or lafora body or lundborg or unverricht) adj2 (disease or syndrome)) or ((jme or jmes) and epilep*) or perioral myoclon*).ti,ab. | | | | | | |
| 11 | infantile spasm/ use emczd, emcr or spasms, infantile/ use ppez or (((early or infantile) adj2 myoclonic adj2 encephalopath*) or ((early or infantile) adj2 epileptic adj2 encephalopath*) or epileptic spasm* or ((flexor or infantile or neonatal) adj2 (seizure* or spasm*)) or generali?ed flexion epileps* or hypsarrhythmia* or ((jacknife or jack nife or lightening or nodding or salaam) adj (attack* or convulsion* or seizure* or spasm*)) or massive myoclonia or minor motor epilepsy or propulsive petit mal or spasm in*1 flexion or spasmus nutans or west syndrome*).ti,ab. | | | | | | |
| 12 | landau kleffner syndrome/ use emczd, emcr, ppez or (dravet or lennox gastaut or lgs or (landau adj2 kleffner) or smei).ti,ab. | | | | | | |
| 13 | lennox gastaut syndrome/ use emczd, emcr or lennox gastaut syndrome/ use ppez or generalized epilepsy/ use emczd, emcr or epileptic syndromes/ use ppez | | | | | | |
| 14 | (child* epileptic encephalopath* or gastaut or lennox or lgs).ti,ab. | | | | | | |
| 15 | myoclonus seizure/ use emczd, emcr or seizures/ use ppez or ((myoclon* adj2 (absence* or epileps* or seizure* or jerk* or progressive familial epilep* or spasm* or convulsion*)) or ((lafora or unverricht) adj2 disease) or muscle jerk).ti,ab. | | | | | | |
| 16 | myoclonic astatic epilepsy/ use emczd, emcr or exp epilepsies, myoclonic/ use ppez or ((myoclonic adj2 | | | | | | |

| (astatic or atonic)) or (myoclonic adj3 (seizure* or spasm*)) or doose* syndrome or mae or generali?ed idiopathic epilepsy).ti,ab. or ((absence or astatic or atonic or tonic or tonic clonic) adj2 (seizure* or spasm*)).ti,ab. 17 exp epilepsies, partial/ use ppez or exp focal epilepsy/ use emczd, emcr or ((focal or focal onset or local or partial or simple partial) adj3 (epileps* or seizure*)).ti,ab. 18 severe myoclonic epilepsy in infancy/ use emczd, emcr or exp epilepsies, myoclonic/ use ppez (dravet*1 or (intractable childhood epilepsy adj2 (generalised tonic clonic or gtc)) or icegtc* or (severe adj2 (myoclonic or polymorphic) adj2 epilepsy adj2 infancy) or smeb or smi).ti,ab. 20 epilepsy, tonic-clonic/ use ppez or epilepsy, generalized/ use ppez or generalized epilepsy/ use emczd, emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* or insult or seizure*))).ti,ab. 21 or/2,4-20 22 exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 22 use ppez 24 budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 25 24 use emczd 26 budget*.ti,ab. 27 cost*.ti. 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 35 limit 34 to engish language | # | searches |
|--|----|--|
| or partial or simple partial) adj3 (epileps* or seizure*)).ti,ab. severe myoclonic epilepsy in infancy/ use emczd, emcr or exp epilepsies, myoclonic/ use ppez (dravet*1 or (intractable childhood epilepsy adj2 (generalised tonic clonic or gtc)) or icegtc* or (severe adj2 (myoclonic or polymorphic) adj2 epilepsy adj2 infancy) or smeb or smei).ti,ab. epilepsy, tonic-clonic/ use ppez or epilepsy, generalized/ use ppez or generalized epilepsy/ use emczd, emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* or insult or seizure*))).ti,ab. or/2,4-20 exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 23 22 use ppez budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ budget*.ti,ab. cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. | | idiopathic epilepsy).ti,ab. or ((absence or astatic or atonic or tonic or tonic clonic) adj2 (seizure* or |
| (dravet*1 or (intractable childhood epilepsy adj2 (generalised tonic clonic or gtc)) or icegtc* or (severe adj2 (myoclonic or polymorphic) adj2 epilepsy adj2 infancy) or smeb or smei).ti,ab. epilepsy, tonic-clonic/ use ppez or epilepsy, generalized/ use ppez or generalized epilepsy/ use emczd, emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* or insult or seizure*))).ti,ab. or/2,4-20 exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 22 use ppez budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 24 use emczd budget*.ti,ab. cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 21 and 33 | 17 | |
| adj2 (myoclonic or polymorphic) adj2 epilepsy adj2 infancy) or smeb or smei).ti,ab. 20 epilepsy, tonic-clonic/ use ppez or epilepsy, generalized/ use ppez or generalized epilepsy/ use emczd, emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* or insult or seizure*))).ti,ab. 21 or/2,4-20 22 exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 23 22 use ppez 24 budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 25 24 use emczd 26 budget*.ti,ab. 27 cost*.ti. 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 18 | severe myoclonic epilepsy in infancy/ use emczd, emcr or exp epilepsies, myoclonic/ use ppez |
| emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* or insult or seizure*))).ti,ab. 21 or/2,4-20 22 exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 23 22 use ppez 24 budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 25 24 use emczd 26 budget*.ti,ab. 27 cost*.ti. 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 19 | |
| exp budgets/ or exp "costs and cost analysis"/ or exp economics, hospital/ or exp economics, medical/ or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 23 22 use ppez 24 budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 25 24 use emczd 26 budget*.ti,ab. 27 cost*.ti. 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 20 | emcr or grand mal epilepsy/ use emczd, emcr or (((clonic or grand mal or tonic or (tonic adj3 clonic)) adj2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* adj (contraction* or convuls* |
| or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or value of life/ 23 22 use ppez 24 budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 25 24 use emczd 26 budget*.ti,ab. 27 cost*.ti. 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 21 | or/2,4-20 |
| budget/ or exp economic evaluation/ or exp fee/ or funding/ or health economics/ or exp health care cost/ 24 use emczd budget*.ti,ab. cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 34 21 and 33 | 22 | or economics, nursing/ or economics, pharmaceutical/ or economics/ or exp "fees and charges"/ or |
| cost/ 24 use emczd budget*.ti,ab. cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 4 21 and 33 | 23 | 22 use ppez |
| budget*.ti,ab. cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 ad 21 and 33 | 24 | |
| cost*.ti. (economic* or pharmaco economic* or pharmacoeconomic*).ti. (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 ad 21 and 33 | 25 | 24 use emczd |
| 28 (economic* or pharmaco economic* or pharmacoeconomic*).ti. 29 (price* or pricing*).ti,ab. 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 26 | budget*.ti,ab. |
| (price* or pricing*).ti,ab. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. (financ* or fee or fees).ti,ab. (value adj2 (money or monetary)).ti,ab. or/23,25-32 ad 21 and 33 | 27 | cost*.ti. |
| 30 (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 28 | (economic* or pharmaco economic* or pharmacoeconomic*).ti. |
| 31 (financ* or fee or fees).ti,ab. 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 29 | (price* or pricing*).ti,ab. |
| 32 (value adj2 (money or monetary)).ti,ab. 33 or/23,25-32 34 21 and 33 | 30 | (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab. |
| 33 or/23,25-32 34 21 and 33 | 31 | (financ* or fee or fees).ti,ab. |
| 34 21 and 33 | 32 | (value adj2 (money or monetary)).ti,ab. |
| | 33 | or/23,25-32 |
| 25 limit 34 to engish language | 34 | 21 and 33 |
| | 25 | limit 34 to engish language |

Database(s): NHS Economic Evaluation Database (NHS EED), HTA database – CRD

Date of last search: 31 March 2021

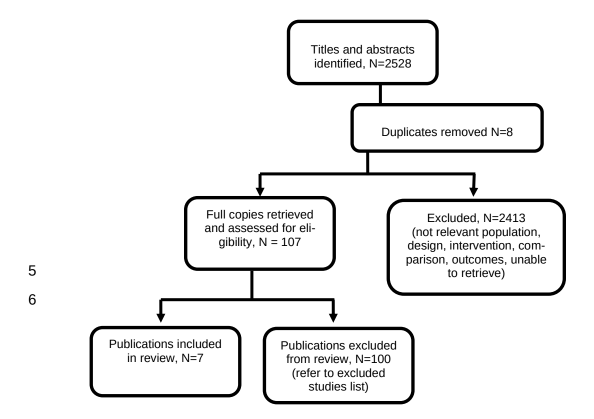
| | Date | e of last search: 31 March 2021 | | | | | | |
|---|------|--|--|--|--|--|--|--|
| | # | searches | | | | | | |
| | 1 | mesh descriptor epilepsy explode all trees | | | | | | |
| | 2 | mesh descriptor seizures this term only | | | | | | |
| | 3 | mesh descriptor seizures, febrile this term only | | | | | | |
| | 4 | mesh descriptor status epilepticus explode all trees | | | | | | |
| | 5 | (epilep* or seizure* or convuls*) or ("continous spike wave of slow sleep" or "infant* spasm*") | | | | | | |
| | 6 | ((absence near2 (convulsion* or seizure*)) or ((typical or atypical) next absenc*) or "petit mal*" or pyknolepsy or "typical absence*") | | | | | | |
| | 7 | mesh descriptor seizures explode all trees | | | | | | |
| | 8 | ((drop or akinetic or atonic or tonic) near2 (attack* or epileps* or seizure* or convulsion*)) or "brief seizure" or (tonic near3 atonic near3 (attack* or epileps* or seizure* or convulsion*)) | | | | | | |
| 9 mesh descriptor epilepsy, rolandic this term only | | | | | | | | |
| | 10 | (bcects or bects or brec or "benign epilepsy" or (benign near2 (childhood or neonatal or pediatric or paediatric) near2 epileps*) or (benign near2 (childhood or neonatal or pediatric or paediatric) near2 (convulsion* or epileps* or seizure* or spasm*)) or (benign near3 (convulsion* or epileps*) near2 centrotemporal near2 spike*) or cects or ((centralopathic or centrotemporal or "temporal-central focal") near (convulsion* or epileps* or seizure*)) or ((osylvian or postrolandic or roland*) near2 (convulsion* or epileps* or seizure* or spasm*))) | | | | | | |
| | 11 | mesh descriptor epilepsy, generalized this term only | | | | | | |
| | 12 | (((akinetic or atonic or central or diffuse or general or generali?ed or idiopathic or tonic) near3 (epilep* or seizure*)) or (("childhood absence" or "juvenile absence" or myoclonic or myoclonia or "myoclonic astatic" or myoclonus or gtcs) near2 epilep*) or (epilepsy near2 "eyelid myoclonia") or (ige near2 phantom absenc*) or "impulsive petit mal" or (janz near3 (epilep* or "petit mal")) or "jeavons syndrome*" or ((janz or lafora or "lafora body" or lundborg or unverricht) near2 (disease or syndrome)) or ((jme or jmes) and epilep*) or "perioral myoclon*") | | | | | | |
| | 13 | mesh descriptor spasms, infantile this term only | | | | | | |
| | 14 | (((early or infantile) near2 myoclonic near2 encephalopath*) or ((early or infantile) near2 epileptic near2 encephalopath*) or "epileptic spasm*" or ((flexor or infantile or neonatal) near2 (seizure* or spasm*)) or "generali?ed flexion epileps*" or hypsarrhythmia* or ((jacknife or "jack nife" or lightening or nodding or salaam) next (attack* or convulsion* or seizure* or spasm*)) or "massive myoclonia" or "minor motor epi- | | | | | | |

| # | searches | | | | | | |
|----|--|--|--|--|--|--|--|
| | lepsy" or "propulsive petit mal" or "spasm in* flexion" or "spasmus nutans" or "west syndrome*") | | | | | | |
| 15 | mesh descriptor landau kleffner syndrome this term only | | | | | | |
| 16 | (dravet or "lennox gastaut" or lgs or (landau near2 kleffner) or smei) | | | | | | |
| 17 | mesh descriptor lennox gastaut syndrome this term only | | | | | | |
| 18 | mesh descriptor epileptic syndromes this term only | | | | | | |
| 19 | ("child* epileptic encephalopath*" or gastaut or lennox or lgs) | | | | | | |
| 20 | ((myoclon* near2 (absence* or epileps* or seizure* or jerk* or "progressive familial epilep*" or spasm* or convulsion*)) or ((lafora or unverricht) near2 disease) or "muscle jerk") | | | | | | |
| 21 | mesh descriptor epilepsies, myoclonic explode all trees | | | | | | |
| 22 | ((myoclonic near2 (astatic or atonic)) or (myoclonic near3 (seizure* or spasm*)) or "doose* syndrome" or mae or "generali?ed idiopathic epilepsy") or ((absence or astatic or atonic or tonic or "tonic clonic") near2 (seizure* or spasm*)) | | | | | | |
| 23 | mesh descriptor epilepsies, partial explode all trees | | | | | | |
| 24 | ((focal or "focal onset" or local or partial or "simple partial") near3 (epileps* or seizure*)) | | | | | | |
| 25 | mesh descriptor epilepsies, myoclonic this term only | | | | | | |
| 26 | (dravet*1 or ("intractable childhood epilepsy" near2 ("generalised tonic clonic" or gtc)) or icegtc* or (severe near2 (myoclonic or polymorphic) near2 epilepsy near2 infancy) or smeb or smei) | | | | | | |
| 27 | mesh descriptor epilepsy, tonic-clonic this term only | | | | | | |
| 28 | mesh descriptor epilepsy, generalized this term only | | | | | | |
| 29 | (((clonic or "grand mal" or tonic or (tonic near3 clonic)) near2 (attack* or contraction* or convuls* or seizure*)) or gtcs or (generali* next (contraction* or convuls* or insult or seizure*))) | | | | | | |
| 30 | #1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 or #22 or #23 or #24 or #25 or #26 or #27 or #28 or #29 | | | | | | |

1 Appendix C - Clinical evidence study selection

- 2 Clinical study selection for review question: What antiseizure therapies
- 3 (monotherapy or add-on) are effective in the treatment of absence seizures?

Figure 1: Study selection flow chart



1 Appendix D – Clinical evidence tables

- 2 Clinical evidence tables for review question: What antiseizure therapies (monotherapy or add-on) are effective in the
- 3 treatment of absence seizures?
- 4 Table 8: Clinical evidence tables

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|----------------------|---------------------------------------|---------------------------|------------------|-------------------------------|-----------------------------|
| Full citation | Sample size | Interventions | Details | Results | Limitations |
| Callaghan, N., | Total recruited: N=28 | Ethosuximide: | Treatment dura- | Primary outcomes | Methodological limita- |
| O'Hare, J., O'Dris- | Ethosuximide group: n=14 | prescribed initially in a | tion: unclear | | tions assessed using the |
| coll, D., Compara- | Sodium valproate group: n=14 | dosage of 250mg per | (mean 3 years, | Remission - complete (no | Cochrane risk of bias |
| tive study of | | day, | range 18 months | longer observed to have at- | tool for randomised trials |
| ethosuximide and | Characteristics | increased by 250mg | - 4 years) | tacks during six-hour te- | (Version 2.0) |
| sodium valproate in | Age, years, mean (range) | increments to a maxi- | | lemetry at two intervals of | |
| the treatment of | Ethosuximide group: 8 (4-14) | mum of 1500mg per | Outcome meas- | six months, and reported to | Domain 1: Randomisa- |
| typical absence sei- | Sodium valproate group: 9 (5-15) | day. Initially prescribed | | have complete freedom from | |
| zures (petit mal), | | to be taken at 1 pm, | hour telemetry | seizures) | 1.1: No information, said |
| Developmental | Gender | but when the dose was | | Ethosuximide group: n=8/14 | to be randomised, not |
| Medicine and Child | Ethosuximide group: 8 male; 6 fe- | increased to more than | every 6 months, | Sodium valproate group: | further information |
| Neurology, 24, | male. | 250mg per day it was | as well as re- | n=6/14 | 1.2: No information |
| 830-836, 1982 | Sodium valproate group: 5 male; 9 | taken in divided doses | ports from par- | | 1.3: Probably no, no dif- |
| | female. | at 8 am and 1 pm. | ents and teacher | Remission - partial (>50% | ferences between base- |
| Ref Id 1080077 | | | | reduction in seizure fre- | line characteristics in the |
| | Age at onset of seizure, years, range | Sodium valproate: | Follow-up: mean | quency during six hour te- | two groups, but very few |
| Country/ies where | Ethosuximide group: 2-5 | prescribed initially in a | 3 years (range | lemetry at two intervals of | baseline characteristics |
| the study was car- | Sodium valproate group: 3-6 | dosage of 400mg dai- | 18 months - 4 | six months, and reported | reported |
| ried out Ireland. | | ly, increased by 200mg | years) | significant reduction in sei- | |
| | Inclusion criteria | increments to a maxi- | | zure frequency) | Domain 2: Deviations |
| Study type RCT. | Patients with absence attacks associ- | mum of 2400mg daily. | | Ethosuximide group: n=3/14 | from intended inter- |
| | ated with three-per-second spike and | Prescribed to be taken | | Sodium valproate group: | ventions: Some con- |
| Aim of the study | wave activity in the EEC | at 8 am and 1 pm. | | n=6/14 | cerns |
| To compare sodium | | | | | 2.1: No information |
| valproate and | Exclusion criteria | | | Remission - none (<50% | 2.2: No information |
| ethosuximide in | Not reported | | | reduction in seizure fre- | 2.3: No information |
| terms of seizure | | | | quency during six hour te- | 2.4: Probably yes, not all |
| control and side- | | | | lemetry at two intervals of | participants had inter- |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|--------------|---------------|---------|--|---|
| effects. Study dates Not reported. Source of funding Labaz, Warner-Lambert Pharmaceuticals. | | | | six months, and reported slight or no improvement) Ethosuximide group: n=3/14 Sodium valproate group: n=2/14* EEG response - normal Ethosuximide group: n=6/14 Sodium valproate group: n=4/14 Adverse events - any Ethosuximide group: n=1/14 Sodium valproate group: n=2/14 | vention as intended 2.5: Probably yes, four participants crossed over due to poor re- sponse to initial drug 2.6: Probably yes Domain 3: Missing outcome data: Low risk 3.1: Yes, data was available for nearly all participants Domain 4: Measure- ment of the outcome: Low risk 4.1: Probably no, out- comes were well defined and assessed by te- lemetry (but not further defined) 4.2: No, outcomes in- cluded seizure frequen- cy and adverse events which are unlikely to differ between treatment arms 4.3: Probably yes, not said to be double blind 4.4: Probably no, out- comes are objective Domain 5: Selection of the reported result: Some concerns 5.1: No information 5.2: No information |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|--|---|--|------------------------------------|---|--|
| | | | | | 5.3: No information |
| | | | | | Domain 6: Overall |
| | | | | | judgement of bi- |
| | | | | | as: Some concerns |
| | | | | | The study is judged to |
| | | | | | raise some concerns in at least one domain, but |
| | | | | | not to be at high risk of |
| | | | | | bias for any domain |
| Full citation | Sample size | Interventions | Details | Results | Limitations |
| Coppola, G., Auricchio, G., Federico, | Total recruited: N=38 Intervention group (Lamotrigine): | Intervention group 0.5 mg/kg/day for 2 | Duration of treatment: not | Primary outcomes | Methodological limita- tions assessed using the |
| R., Carotenuto, M., | n=19 | weeks in two divided | reported. | Seizure free, n (no clinical | Cochrane risk of bias |
| Pascotto, A., | Control group (Valproic Acid): n=19 | doses, followed by 1.0 | . op o. to d. | absences reported by exter- | tool for randomised trials |
| Lamotrigine versus | | mg/kg/day for an addi- | Outcome meas- | nal observers for at least the | (Version 2.0) |
| valproic acid as | Characteristics | tional 2 weeks. There- | urement: Pa- | previous month, and no | Damain 4. Dandamiaa |
| first-line monother- apy in newly diag- | Age, years, mean (range) Total sample: 7.5 (3-13) | after, doses were in- creased in 1- | tients were seen at monthly inter- | electroclinical seizures de- tected in awake video-EEG | Domain 1: Randomisa- tion: Low risk |
| nosed typical ab- | Total Sample. 7.5 (5-15) | mg/kg/day increments | vals for ≤12 | with HV-EEG and in 24-h | 1.1: No information, just |
| sence seizures: An | Age at seizure onset, years, mean | every 5 days until sei- | months, and | ambulatory EEG monitoring) | said to be randomised |
| open-label, random- | | zures were controlled | were questioned | , | 1.2: Probably yes, |
| ized, parallel-group | Intervention: 7.5 (4-12) | (as indicated by lack of | | 1 month | makes reference to an |
| study, Epilepsia, 45, | Control: 7.5 (3-13) | clinical evidence of | fects (recorded | Intervention group 1/19 | external investigator |
| 1049-1053, 2004 | Males, n (%) | absences and no elec- troclinical seizures in | in a diary), a medical exami- | Control group 10/19 | 1.3: No, no significant differences between |
| Ref Id | Intervention: 7 (36.8) | an awake video-EEG | nation and a vid- | 3 months | groups at baseline |
| 1080163. | Control: 10 (52.6) | with HVEEG and in a | eo EEG record- | Intervention group: n=7/19 | |
| | | 24-h ambulatory EEG), | | Control group: n=12/19 | Domain 2: Deviations |
| Country/ies where | Inclusion criteria | intolerable adverse | ed HV-IPS. A 24 | 40 | from intended inter- |
| the study was car- ried out | Age from 3 to 13 years | effects occurred, or a maximum dose of 12 | hour ambulatory EEG was per- | 12 months Intervention group: n=10/19 | ventions: Low risk 2.1: Yes, open label |
| Italy | Newly diagnosed typical absence | mg/kg/day had been | formed if there | Control group: n=13/19 | study |
| , | seizures associated with general- | reached. the maximum | | 5.05p711 10,10 | 2.2: Yes, open label |
| Study type | ized, synchronous 3-Hz (2.5–4 Hz) spike-and-wave activity, lasting >3 | allowed dose of LTG in | of absences. | Adverse events - any, n | study |
| RCT | s, occurring spontaneously or dur- | patients completing | Data analysed | (%) number of participants | |
| Aim of the study | ing one of two trials of 3-min hyper- | without interruption the full uptitration schedule | | experiencing an adverse event | Domain 3: Missing outcome data: Low |
| i or the study | | ian aparation concadio | tornion to trout | 0.011 | Jaioonio datai Eon |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|--|--|---|---|--|--|
| To compare the efficacy of lamotrigine and valproic acid in newly diagnosed children and adolescents with typical absence seizures Study dates Not reported (publication date 2004) Source of funding Not reported | ventilation with a 1- to 2-min rest between trials Clearly observable clinical signs of typical absence seizures (for example, staring or impairment of consciousness) on the video record Normal clinical, neurologic, and computed tomography (CT)/magnetic resonance imaging (MRI) examination Informed consent by parents or caregivers Exclusion criteria Absences with marked eyelid or perioral myoclonia with absences) Absences with marked limb and trunk rhythmic myoclonic jerks (myoclonic absence epilepsy) Absences with single ictal myoclonic jerks of the limbs, trunk, or head Absences with mild or not clinically detectable impairment of consciousness (for example, juvenile myoclonic epilepsy) Other types of epileptic seizures Stimulus-sensitive absences: photosensitive, patternsensitive, self-induced pattern-sensitive Irregular, arrhythmic spike/multiple spike and slow-wave EEG discharges with marked variations of discharge frequency Central-temporal or occipital focal | was reached ~75 days after initiation of treatment. Control group Administered as 200-mg enteric-coated non–sustained-release sodium valproate tablets or as liquid formulation (40 mg/ml), started at 10 mg/kg/day and increased by 5 mg/kg/day every 3 days until seizures were controlled or intolerable side effects occurred, up to a maximum of 30 mg/kg/day given in three divided doses. | Follow-up: 12 months (assessements took place at 1 month, 3 months, and 12 months). | Intervention group: n=6/19 Control group: n=2/19 Treatment cessation due to adverse events, n number of participants withdrawing from treatment due to adverse events Intervention group: n=0/19 Control group: n=0/19 Withdrawal from study (by 3 months) Intervention group n=6/19 Control group n=3/19. Withdrawal from study (by 12 months) Intervention group n=6/19 Control group n=3/19. | risk 3.1: Yes, data was available for all participants randomised Domain 4: Measurement of the outcome: Low risk 4.1: No, outcomes were well defined and was assessed using reliable method (EEG) 4.2: No, outcomes included seizure free status and side effects, which are unlikely to differ between treatment arms 4.3: No, said to be unaware of treatment allocation Domain 5: Selection of the reported result: Some concerns 5.1: No information, there is no reference to any study protocol 5.2: No information. Trial protocol was not available 5.3: No information. Trial protocol was not available Domain 6: Overall judgment of bias: Some concerns |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|---|---|-----------------|---|---|
| | EEG discharges or abnormal background EEG activity Known or suspected structural brain lesion Progressive neurologic illness Psychiatric disorder requiring medication Chronic cardiovascular, renal, or hepatic disease and, in general, any disease that could interfere with drug absorption, distribution, metabolism, or excretion Long-term comedication with other drugs Suspected poor compliance | | | | The study is judged to raise some concerns in at least one domain, but not to be at high risk of bias for any domain |
| Full citation Fattore, C., Boniver, C., Capovilla, G., Cerminara, C., Cit- terio, A., Coppola, G., Costa, P., Dar- ra, F., Vecchi, M., Perucca, E., A mul- ticenter, random- ized, placebo- controlled trial of levetiracetam in children and ado- lescents with newly diagnosed absence epilepsy, Epilepsia, 52, 802-809, 2011 Ref Id 1080326 | Sample size Total recruited: N=59 Intervention group (levetiracetam): n=38 Control group (placebo): n=21 Characteristics Age, years, mean (SD, range) Intervention: 8.7 (2.2, 4.9 to 13.0) Control: 7.9 (3.0, 4.0 to 15.0) Males, n Intervention: 15 Control: 12 Syndromic diagnosis, childhood absence epilepsy (n) Intervention: 34 Control: 20 Syndromic diagnosis, juvenile absence epilepsy (n) Intervention: 4 Control: 1 | lowed by 15 mg/kg/day for the next 4 days (days 4–7). If tolerabil- | recordings were | Results Primary outcomes Responder status (no clinical seizures on days 13 and 14, and no EEG seizures during the standard EEG on day - ITT analysis Levetiracetam n=9/38 Placebo n=1/21 Patients free from clinical and EEG seizures on days 4–7 (this is, first week of treatment when levetiracetam dose could not exceed 15 mg/kg/day) Intervention group: n=4/38 Control group: n=0/21 | Limitations Methodological limitations assessed using the Cochrane risk of bias tool for randomised trials (Version 2.0) Domain 1: Randomisation: Low risk 1.1: Yes, randomised using computergenerated random numbers 1.2: Probably yes, makes reference to maintaining blinding for subsequent assignments which were disclosed at study end 1.3: Probably no, some differences for example |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|--|---|---------|--|--|
| Country/ies where the study was carried out Italy Study type Multi-centre, randomised, placebocontrolled trial Aim of the study To evaluate the potential efficacy of levetiracetam as an antiabsence agent in children and adolescents with newly diagnosed childhood or juvenile absence epilepsy Study dates Patient enrolment and treatment: October 2006- | Seizure frequency at pretreatment, >10/day Intervention: 26 Control: 18 Seizure frequency at pretreatment, 6- 10/day Intervention: 6 Control: 0 Seizure frequency at pretreatment, 1- 5/day Intervention: 4 Control: 2 Seizure frequency at pretreatment, 1- 6/week Intervention: 2 Control: 1 Inclusion criteria Age between 4 and 16 years Recent diagnosis of childhood or juvenile absence epilepsy, as defined by the International League Against Epilepsy (ILAE) criteria (Commission, 1989) Electroencephalographic (EEG) evidence of regular, synchronous and symmetrical spike-wave paroxysmal discharges with a frequency of about 3 Hz and duration of at least 4 s, occurring spontaneously or during hyperventilation (a 2–3-min hyper-ventilation test, followed 2 min later by a second hyperventilation if paroxysms were not detected in the first test) or intermittent photic stimulation (IPS; 5 s stimulation at each frequency in the 10–30 Hz | 30 mg/kg/day on day 11 and maintained until day 14. If no clini- cal or EEG seizures | Methods | and EEG seizures on days 11–14 Intervention group: n=7/38 Control group: n=0/21 50% reduction (vs baseline) in total duration of EEG seizures on day 14 Intervention group: n=12/38 Control group: n=3/21 Adverse events - any Intervention group: n=7/38 Control group: n=3/21 Adverse events - serious Intervention group n=0/38 Control group n=0/21 Adverse events - thought to be related to treatment Intervention group: n=3/38 Control group: n=0/21 Adverse events - leading to discontinuation Intervention group: n=1/38 Control group: n=0/21 | gender, but not significant Domain 2: Deviations from intended interventions: Low risk 2.1: No, said to be double blind 2.2: No, said to be double blind 2.3: N/A 2.4: Probably no, no mention of deviations from the intervention apart from one withdrawal 2.5: Probably no, nearly all participants received the intervention 2.6: N/A Domain 3: Missing outcome data: Low risk 3.1: Yes, data was available for all but 1 participants randomised Domain 4: Measurement of the outcome: Low risk 4.1: No, outcomes were well defined and was assessed using reliable method (EEG) 4.2: No, outcomes included seizure free status and side ef- |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---------------|---|--|---------|----------------------|---|
| | range, eye closed) A history of clinically evident spontaneously occurring absence seizures impacting on functional abilities Written informed consent. Exclusion criteria A history of generalized tonic clonic seizures Clinical or EEG findings inconsistent with a diagnosis of childhood or juvenile absence epilepsy Previous treatment with ASMs (except for earlier treatments for other indications such as febrile seizures, or brief exposures to other ASMs prior to diagnostic assessment) Intellectual disability Clinically significant hepatic or renal disorders History of hypersensitivity reactions to study products or structurally related substances Any condition that, in the investigator's judgment, was expected to impact negatively on subjects' health or study procedures. | body weight, and preference, either a liquid formulation (100 mg/ml solution) or 500 mg tablets were used, with matching placebos. Both treatments were administered in the morning and in the evening in two equally divided doses | | | fects, which are unlikely to differ between treatment arms 4.3: No, randomisation was only broken after outcome measure evaluation Domain 5: Selection of the reported result: Some concerns 5.1: No information, there is no reference to any study protocol 5.2: No information. Trial protocol was not available 5.3: No information. Trial protocol was not available Domain 6: Overall judgement of bias: Some concerns The study is judged to raise some concerns The study is judged to raise some concerns in at least one domain, but not to be at high risk of bias for any domain Other information After the 2 week double-blind period, all placebo participants were started on levetiracetam. Some participants in both groups discontinued with levetiracetam and start- |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|--|---|---|---|---|
| | | | | | ed other drugs, for example, valproate. |
| T., Holmes, G. L., Manasco, P., Concannon, S., Chen, C., Womble, G., Casale, E. J., Lamictal (lamotrigine) monotherapy for typical absence seizures in children, Epilepsia, 40, 973- 979, 1999 Ref Id 1080361 Country/ies where the study was carried out USA Study type Double blind, placebo controlled study Aim of the study To investigate | Sample size Total recruited and took part in dose escalation phase: N=45. Total randomised after dose escalation phase: Lamotrigine group: n=15 Placebo group: n=14 Total analysed Lamotrigine group: n=14 (1 patient in the lamotrigine, group withdrew consent) Placebo group: n=14 (1 patient failed to meet the 80% compliance standard; however they were included in the analysis on an intent-to-treat basis). Characteristics Age, years, mean (SD) Lamotrigine group: 6.9 (2.3) Placebo group: 8.8 (3.1) Gender Lamotrigine group: 36% males, 64% females Placebo group: 36% males, 64% females Placebo group: 36% males, 64% females Inclusion criteria Patients 2-16 years old with newly diagnosed typical absence seizures, as evidenced by the clinical and EEG features of typical absence seizures on one of two 3-min | Interventions Before randomisation to lamotrigine or placebo, all participants underwent a dose escalation phase to identify the dose at which they would be rendered seizure free. Lamotrigine: Initial dose of 0.5mg/kg/day for 2 weeks, followed by 1mg/kg/day for 2 weeks. Following increases by 1mg/kg/day according to response, maximum dose of 15mg/kg/day. Chewable 5mg, 25 and 100mg tables were provided. Continued at the dose identified during the dose escalation stage for 4 weeks. No adjustment of dose was permitted. Placebo: The dose of lamotrigine during the dose escalation phase was tapered to placebo over 2 weeks: patients received 50, 25%, and 0 (this is, | of dose escalation, and end of the RCT phase. Follow-up: 4 weeks (RCT study only). | Results Primary outcomes Remained seizure free at end of placebo controlled phase Lamotrigine group: 9/14 Placebo group: 3/14 Adverse events - leading to withdrawal from study Lamotrigine group: n=0/14 Placebo group: n=0/14 | Limitations Methodological limitations assessed using the Cochrane risk of bias tool for randomised trials (Version 2.0) Domain 1: Randomisation: Some concerns 1.1: No information, said to be randomised, not further information 1.2: No information 1.3: No, no significant differences at baseline between groups Domain 2: Deviations from intended interventions: Low risk 2.1: No, said to be double blind 2.2: No, said to be double blind 2.3: N/A 2.4: Probably no, no mention of deviations from the intervention 2.5: Probably no, nearly all participants received the intervention apart from one participant who was not compliant 2.6: N/A |
| | hyperventilation tests | 100% placebo) of their | | | Domain 3: Missing |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|--|--|---------------|---------|----------------------|---|
| Study dates Not reported Source of funding Glaxo Wellcome Inc. (Research Tri- angle Park, NC) | Any woman of reproductive capability was required to use a contraceptive method acceptable to the investigator and to provide a written statement of intent to avoid pregnancy Exclusion criteria A known or suspected structural lesion A history of poor compliance with medication or abuse of drugs A progressive neurologic illness (defined prospectively as being an unstable pathologic state during the previous 12 months) A psychiatric disorder requiring medication Chronic cardiovascular, renal, or hepatic disease Use of an investigational drug within the previous 12 weeks Any disease thought to interfere with absorption, distribution, metabolism, or excretion of drugs in general With the exception of patients taking methylphenidate, dexamphetamine, or clonidine to treat attention-deficit hyperactivity disorder, patients were not allowed to take psychotropic drugs at any time during the study. | weeks. | | | outcome data: Low risk 3.1: Yes, data was available for all but 1 participants randomised Domain 4: Measurement of the outcome: Low risk 4.1: No, outcomes were well defined and was assessed using reliable method (EEG) 4.2: No, outcomes included seizure free status and side effects, which are unlikely to differ between treatment arms 4.3: Probably no, EEG recordings were analysed by an investigator who was unaware of treatments or doses Domain 5: Selection of the reported result: Some concerns 5.1: Probably yes, there is a protocol however it is not detailed 5.2: Probably no Domain 6: Overall judgement of bias: Some concerns The study is judged to raise some concerns in |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|---|---|--------------------------------|---|---|
| | | | | | at least one domain, but not to be at high risk of bias for any domain. Other information 28 patients completed double-blind phase phase (one patient in the LTG, group withdrew consent). Only one patient (in the placebo- |
| | | | | | treated group) failed to meet the 80% compli- ance standard; this pa- tient was included in analysis on an intent-to- treat basis. |
| Full citation Glauser, T. A., | Sample size Total randomised: N=453 | Interventions Ethosuximide (Zaron- | Details Treatment dura- | Results Primary outcomes | Limitations Methodological limita- |
| Cnaan, A., Shinnar, | Ethosuximide group: n=156 | tin) | tion: 12 months. | Primary outcomes | tions assessed using the |
| S., Hirtz, D. G., | Lamotrigine group: n=149 | 250-mg capsules or | | Freedom from treatment | Cochrane risk of bias |
| Dlugos, D., Masur, | Valproic acid group: n=148 | 250 mg per 5 ml of | Outcome meas- | failure at 16 or 20 weeks | tool for randomised trials |
| D., Clark, P. O., Capparelli, E. V., | Characteristics | syrup. The highest allowable daily dose was | urement: seizure | (treatment failure defined as persistence of absence | (Version 2.0) |
| Adamson, P. C., | Age ≥6 years, n (%) | • | termined by clin- | seizures, a generalised ton- | Domain 1: |
| Ethosuximide, | Ethosuximide group: 116 (75) | body weight, to a max- | | ic-clonic seizure at any time, | |
| valproic acid, and | Lamotrigine group: 110 (74) | | side hyperventi- | excessive drug-related sys- | concerns |
| lamotrigine in child- | Valproic acid group: 113 (77) | | lation testing, | temic toxicity of at least 3.0 | 1.1: Yes, said to be |
| hood absence epi- | Male sex, n (%) | Lamotrigine (Lamictal) | | from baseline, dose-limiting | computer generated |
| lepsy, New England | Ethosuximide group: 65 (42) Lamotrigine group: 57 (38) | 5-mg and 25-mg chewable tablets or | EEG | toxicity after a single downward dose modification, and | 1.2: Probably yes, treatment assignments |
| | Valproic acid group: 71 (48) | 25-mg tablets. The | Data analysed | withdrawal initiated by the | were performed centrally |
| | | highest allowable daily | according | parent or physician): | 1.3: No, no significant |
| Ref Id | No statistically significant differences | dose was 12 mg per | to intention-to- | | differences at baseline |
| 1082357 | seen between the treatment groups | kilogram of body weight, to a maximum | treat approach | Ethosuximide: 81/154 Lamotrigine: 43/146 | between groups |
| Country/ies where | | of 600 mg/day. | Follow-up: 12 | Valproic acid: 85/146 | Domain 2: Deviations |
| the study was car- | • Between 2.5 and 13 years of age | | months. This | | from intended inter- |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|--|---|--|---|---|
| ried out USA Study type Multicentre, double- blind, randomised controlled trial Aim of the study To assess the efficacy, tolerability, and neuropsychological effect of ethosuximide, valproic acid, and lamotrigine to determine the optimal initial empirical monotherapy for children with childhood absence epilepsy Study dates July 2004 to October 2007 Source of funding Supported by grants from the National Institutes of Health | Childhood absence epilepsy of new onset that was clinically diagnosed according to the International League Against Epilepsy classification of epilepsy syndromes (including frequent clinical absence seizures and reported normal development) Had bilateral synchronous, symmetric spike waves (2.7 to 5 Hz) on a normal background with at least one electrographically recorded seizure lasting 3 seconds or more on a 1-hour, awake video EEG Weighed 10 kg or more Had a body-mass index below the 99th percentile Had a normal complete blood count and normal levels of serum alanine aminotransferase, serum aspartate aminotransferase, and bilirubin The girls had to be premenarchal Exclusion criteria Had received antiseizure medication for more than 7 days before randomization Had a history of nonfebrile seizures other than absence seizures (for example, afebrile generalized tonic—clonic or myoclonic seizures) Had a history consistent with juvenile absence epilepsy or juvenile myoclonic epilepsy (for example, generalized tonic—clonic or myoclonic seizures) | body weight, to a maximum of 3000 mg/day. | findings at 16 or 20 weeks (prima- ry outcome data based on as- sessments at week 16 unless unless the partic- ipant had a 5th visit at 20 weeks, in which case data from the 20 | Ethosuximide: 37/154 Lamotrigine: 25/146 Valproic acid: 35/146 Study drug discontinued – | 2.1: Probably no, said to be double blind 2.2: Probably no, said to be double blind 2.3: N/A 2.4: N/A 2.5: N/A 2.6: Yes Domain 3: Missing outcome data: Low risk 3.1: Yes, data was available for nearly all participants Domain 4: Measurement of the outcome: Low risk 4.1: No, outcomes were well defined and was assessed using reliable method (EEG) 4.2: No, outcomes included seizure free control and side effects, which are unlikely to differ between treatment arms 4.3: No information |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|--|--|------------------|---------------------|--|---|
| | Had a history of a severe dermatologic reaction to any medication Had a history of major psychiatric disease, autistic-spectrum disorder, or any clinically significant medical condition | | | | 5.1: Probably yes, there is a protocol however it is not detailed 5.2: Probably no 5.3: Probably no Domain 6: Overall judgement of bias: Low The study is judged to be at low risk of bia |
| Full citation Glauser TA, Cnaan A, Shinnar S, Hirtz DG, Dlugos D, Masur D, Clark PO, Adamson PC, Childhood Absence Epilepsy Study Team. Ethosuximide, valproic acid, and lamotrigine in childhood absence epilepsy: initial monotherapy outcomes at 12 months. Epilepsia. 2013 Jan;54(1):141-55. | See Glauser 2010 | See Glauser 2010 | See Glauser 2010 | Primary outcomes Freedom from treatment failure at 12 months (treatment failure defined as persistence of absence seizures, a generalised ton- ic-clonic seizure at any time, excessive drug-related sys- temic toxicity of at least 3.0 from baseline, dose-limiting toxicity after a single down- ward dose modification, and withdrawal initiated by the parent or physician): Ethosuximide: 70/154 Lamotrigine: 31/146 Valproic acid: 64/146 Adverse events - serious (12 months) Ethosuximide: 4/155 Lamotrigine: 2/149 Valproic acid: 2/147 Intolerable adverse effects (12 months) Ethosuximide: 38/154 Lamotrigine: 29/146 | See Glauser 2010 |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|---|---|--|--|---|
| | | | | Valproic acid: 48/146 Withdrawal from study (12 months) Ethosuximide: 29/154 Lamotrigine: 18/146 Valproic acid: 21/146 Attentional dysfunction defined as a Confidence Index of 0.60 or higher on the Conners' Continuous Performance Test; 12 months): Ethosuximide: 20/70 Lamotrigine: 8/30 Valproic acid: 34/61 | |
| ethosuximide in the treatment of ab- sence seizures, Neurology, 32, 157- 163, 1982 Ref Id 1115033 | Valproic acid during period I group: n=22 (treatment naïve n=7, refractory | Interventions Valproic acid and placebo Ethosuximide (followed by Ethosuximide and placebo Valproic acid in period II): For group 1 (first 23 patients enrolled): Daily dosage of Valproic acid started at 15-20 mg per kg and 5 days later increased to a maximum of 30 mg/kg if the 12-hour EEG showed generalised spike wave discharges For group 2 (the next 22 patients enrolled): Initial daily dosage of 12.5 to 20 mg/kg, with a dosage increase | Details Duration of treatment: 6 weeks of period I drug, followed by 6 weeks of peri- od II drug Outcome meas- urement: as- sessment of sei- zure control took place at 6 and 12 weeks of treatment. Follow-up: 6 weeks (period 1 only). | naive patients (100% reduction in spike wave bursts on EEG) Ethosuximide in period 1 n=4/9. Valproic acid in period 1: n=6/7 | Cochrane risk of bias tool for randomised trials (Version 2.0) Domain 1: Randomisation: Some concerns 1.1: No information, said to be randomly as- |

| Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|---|---|--|---------|----------------------|--|
| Aim of the study To evaluate the efficacy of VPA in treating absence seizures Study dates Not reported Source of funding The NINCDS | been observed by one of the investigators and must have occurred at least once on the pretreatment 12-hour telemetered EEG. Some patients also had other types of seizures. No evidence of progressive neurologic illness must have been kept on the maximally tolerated daily dosage of ESM for 1 month before the study Exclusion criteria Not reported | every 2 days for 2 weeks, to a maximum of 60 mg/kg/day Ethosuximide and Valproic acid placebo (followed by Valproic acid and Ethosuximide placebo in period II): Group 1: Daily dosage ranged from 250 to 1500mg Group 2: Daily dosage was 250 to 1500 mg. For both groups anti- seizure drugs for the treatment of other sei- zure types were con- tinued throughout the study. | | | 2.1: Probably no, said to be double blind but no details 2.2: Probably no, said to be double blind but no details 2.3: N/A 2.4: Probably no, no mention of deviations from the intervention 2.5: Probably no, for period I there were no drop outs. Some participants did not cross over to period II 2.6: N/A Domain 3: Missing outcome data: Low risk 3.1: Yes, data was available for nearly all participants for period I Domain 4: Measurement of the outcome: Low risk 4.1: No, outcomes were well defined and was assessed using reliable method (EEG) 4.2: No, outcomes included seizure frequency which is unlikely to differ between treatment arms 4.3: No information 4.4: Probably no, out- |

| comes are objective Domain 5: Selection of the reported result: Some concerns 5.1: No information 5.2: No information 5.3: No information 5.3: No information Domain 6: Overall judgement of bias: Some concerns The study is judged to raise some concerns in at least one domain, but not to be at high risk of bias for any domain Other information Both naive and refractory patients were enrolled. Naive patients who had 100% seizure control in the first phase, and refractory patients who had at least 80% seizure control in the first phase were not crossed over to a different drug in phase II, but | Study details | Participants | Interventions | Methods | Outcomes and Results | Comments |
|--|---------------|--------------|---------------|---------|----------------------|---|
| instead maintained on the same drug. Only those who did not re- | Study details | Participants | Interventions | Methods | Outcomes and Results | Domain 5: Selection of the reported result: Some concerns 5.1: No information 5.2: No information 5.3: No information Domain 6: Overall judgement of bias: Some concerns The study is judged to raise some concerns in at least one domain, but not to be at high risk of bias for any domain Other information Both naive and refractory patients were enrolled. Naive patients who had 100% seizure control in the first phase, and refractory patients who had at least 80% seizure control in the first phase were not crossed over to a different drug in phase II, but instead maintained on the same drug. Only |

¹ EEG: electroencephalograml; ESM: ethosuximide; LTG: lamotrigine; RCT: randomised controlled trial;SD: standard deviation; VPA: valproic acid

1 Appendix E – Forest plots

- 2 Forest plots for review question: What antiseizure therapies (monotherapy or
- add-on) are effective in the treatment of absence seizures?
- 4 Comparison 2: lamotrigine versus valproic acid
- 5 Figure 2: withdrawal from study (12 months)



6 7

1 Appendix F – GRADE tables

- 2 GRADE tables for review question: What antiseizure therapies (monotherapy or add-on) are effective in the treatment of
- 3 absence seizures?
- 4 Table 9: Clinical evidence profile. Comparison 1: ethosuximide versus sodium valproate

| Quality ass | essment | | | | Assessment Number of patients Effect | | | Quality | Importance | | | |
|---------------------|--------------|-------------------------------|-----------------------------|----------------------------|--------------------------------------|----------------------|-------------------|-------------------|------------------------------|---|------------------|----------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ETH | VPA | Relative (95% CI) | Absolute | | |
| Seizure free | e at 6 weeks | s - naive | | | | | | | | | | |
| 1 (Sato 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | serious ² | none | 4/9 (44.4%) | 6/7 (85.7%) | RR 0.52 (0.24 to 1.14) | 411 fewer per 1000 (from 651 fewer to 120 more) | ⊕⊕OO LOW | CRITICAL |
| Seizure free | e at 6 weeks | s - refractory | | | | | | | | | | |
| 1 (Sato 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 4/14 (28.6%) | 3/15 (20%) | RR 1.43 (0.39 to 5.28) | 86 more per 1000 (from 122 fewer to 856 more) | ⊕OOO VERY LOW | CRITICAL |
| Freedom fre | om treatme | nt failure at 1 | 6 or 20 weeks* | | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 81/154 (52.6%) | 85/146 (58.2%) | RR 0.9 (0.74 to 1.11) | 58 fewer per 1000 (from 151 fewer to 64 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Freedom fre | om treatme | nt failure at 1 | 2 months* | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 70/154 (45.5%) | 64/146 (43.8%) | RR 1.04 (0.81 to 1.33) | 18 more per 1000 (from 83 fewer to 145 more) | ⊕⊕⊕O MODERATE | CRITICAL |

| Quality ass | uality assessment | | | | | Number of patients Effect | | | Quality | Importance | | |
|----------------------------|--|-------------------------------|-----------------------------|----------------------------|---------------------------|---------------------------|-----------------|-----------------|------------------------------|---|------------------|----------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ETH | VPA | Relative (95% CI) | Absolute | | |
| 1 (Calla- ghan 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 8/14 (57.1%) | 6/14 (42.9%) | RR 1.33 (0.63 to 2.84) | 141 more per 1000 (from 159 fewer to 789 more) | ⊕000 VERY LOW | CRITICAL |
| Remission | - partial (>5 | 0% reduction | in seizure freque | ency during six h | our telemetry a | t two intervals of s | ix months, | and reporte | ed significant re | eduction in seiz | zure frequency) | |
| 1 (Calla- ghan 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 3/14 (21.4%) | 6/14 (42.9%) | RR 0.5 (0.15 to 1.61) | 214 fewer per 1000 (from 364 fewer to 261 more) | ⊕OOO VERY LOW | CRITICAL |
| Remission | Remission - none (<50% reduction in seizure frequency during six hour telemetry at two intervals of six months, and reported slight or no improvement) | | | | | | | | | | | |
| 1 (Calla- ghan 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 3/14 (21.4%) | 2/14 (14.3%) | RR 1.5 (0.29 to 7.65) | 71 more per 1000 (from 101 fewer to 950 more) | ⊕000 VERY LOW | CRITICAL |
| EEG respoi | nse - norma | ıl (follow-up 1 | 8 months - 4 year | rs. mean = 3 vear | ·s) | | | | | | | |
| 1 (Calla- ghan 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 6/14 (42.9%) | 4/14 (28.6%) | RR 1.5 (0.54 to 4.18) | 143 more per 1000 (from 131 fewer to 909 more) | ⊕000 VERY LOW | CRITICAL |
| Adverse ev | ents - anv (| follow-up 18 | months - 4 years, | mean = 3 years) | | | | | | | | |
| 1 (Calla- ghan 1982) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 1/14 (7.1%) | 2/14 (14.3%) | RR 0.5 (0.05 to 4.9) | 71 fewer per 1000 (from 136 fewer to 557 more) | ⊕000 VERY LOW | CRITICAL |
| Adverse ev | ents - serio | us (follow-un | 16 or 20 weeks) | | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 4/155 (2.6%) | 2/147 (1.4%) | RR 1.9 (0.35 to 10.2) | 12 more per 1000 (from 9 fewer to | ⊕⊕OO LOW | CRITICAL |

| Quality ass | essment | | | | | | Number | of patients | Effect | | Quality | Importance |
|---------------------|--------------|-------------------------------|-----------------------------|----------------------------|---------------------------|----------------------|-------------------|-------------------|------------------------------|---|------------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ЕТН | VPA | Relative (95% CI) | Absolute | | |
| | | | | | | | | | | 125 more) | | |
| Adverse ev | ents - serio | us (follow-up | 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 4/155 (2.6%) | 2/147 (1.4%) | RR 1.9 (0.35 to 10.2) | 12 more per 1000 (from 9 fewer to 125 more) | ⊕⊕OO LOW | CRITICAL |
| Intolerable | adverse eff | ects (follow-u | ıp 16 or 20 weeks |) | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 37/154 (24%) | 35/146 (24%) | RR 1 (0.67 to 1.5) | 0 fewer per 1000 (from 79 fewer to 120 more) | ⊕⊕OO LOW | CRITICAL |
| Intolerable | adverse eff | ects (follow-u | ip 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 38/154 (24.7%) | 48/146 (32.9%) | RR 0.75 (0.52 to 1.08) | 82 fewer per 1000 (from 158 fewer to 26 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Study drug | discontinu | ed – no reaso | on reported (follow | w-up 16 or 20 we | eks) | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ⁴ | none | 0/154 (0%) | 0/146 (0%) | RD 0.00 (- 0.01 to 0.01) | 0 more per 1000 (from 10 fewer to 10 more) | ⊕⊕OO LOW | CRITICAL |
| Withdrawal | from study | (follow-up 16 | 6 or 20 weeks) | | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 20/154 (13%) | 15/146 (10.3%) | RR 1.26 (0.67 to 2.37) | 27 more per 1000 (from 34 fewer to 141 more) | ⊕⊕OO LOW | CRITICAL |
| Withdrawal | from study | (follow-up 12 | 2 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 29/154 (18.8%) | 21/146 (14.4%) | RR 1.31 (0.78 to 2.19) | 45 more per 1000 (from 32 fewer to | ⊕⊕OO LOW | CRITICAL |

| Quality ass | sessment | | | | | | Number | of patients | Effect | | Quality | Importance |
|---------------------|-------------|-------------------------------|-----------------------------|-------------------------------|---------------------------|----------------------|------------------|-------------------|------------------------------|--|------------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ETH | VPA | Relative (95% CI) | Absolute | | |
| | | | | | | | | | | 171 more) | | |
| Conners' C | ontinuous F | Performance | Test score > 0.60 | [¥] (follow-up 16 or | 20 weeks) | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 35/106 (33%) | 52/106 (49.1%) | RR 0.67 (0.48 to 0.94) | 162 fewer per 1000 (from 29 fewer to 255 fewer) | ⊕⊕⊕O MODERATE | IMPORTANT |
| Conners' C | ontinuous F | Performance | Test score > 0.60 | [¥] (follow-up 12 m | onths) | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 20/70 (28.6%) | 34/61 (55.7%) | RR 0.51 (0.33 to 0.79) | 273 fewer per 1000 (from 117 fewer to 373 fewer) | ⊕⊕⊕⊕ HIGH | IMPORTANT |

^{*} Treatment failure defined as persistence of absence seizures, a generalised tonic-clonic seizure at any time, excessive drug-related systemic toxicity of at least 3.0 from baseline, dose-limiting toxicity after a single downward dose modification, and withdrawal initiated by the parent or physician.

*An index of 0.60 corresponds to a 60% probability that the child has clinically significant attention deficit disorder

1 Serious risk of bias in the evidence contributing to the outcomes as per RoB 2

Table 10: Clinical evidence profile. Comparison 2: lamotrigine versus sodium valproate

| 1 (Coppolar RCT serious no serious none 1/19 10/19 RR 0.1 474 fewer per 10/19 (5.3%) (52.6%) (0.01 to 1000 (from 153 fewer to 521) | Quality ass | essment | | | | | | Number o | of patients | Effect | | Quality | Importance |
|---|--------------|---------------------------|----------------------|---------------|--------------|-------------|------|----------|-------------|----------|----------------|---------|------------|
| 1 (Coppolar 2004) RCT serious¹ no serious inconsistency indirectness imprecision no serious no serious indirectness imprecision no serious no serious indirectness imprecision 1/19 10/19 RR 0.1 474 fewer per 1000 (from 153 fewer to 521 MODERATE | | Design | | Inconsistency | Indirectness | Imprecision | | LTG | VPA | | Absolute | | |
| 1 (Coppo- RCT serious no serious no serious no serious no serious no serious none 1/19 10/19 RR 0.1 474 fewer per head of the final serious none inconsistency indirectness imprecision (5.3%) (52.6%) (0.01 to 1000 (from 153 fewer to 521) | Seizure free | Seizure freedom - 1 month | | | | | | | | | | | |
| rewer) | | RCT | serious ¹ | | | | none | | | (0.01 to | 1000 (from 153 | | CRITICAL |

Confidence interval crosses one MID (0.8 or 1.25)
 Confidence intervals cross both MIDs (0.8 and 1.25)
 Absolute effect range crosses 2 MIDs (10 more per 1000 and 10 fewer per 1000)

| Quality asso | essment | | | | | | Number o | of patients | Effect | | Quality | Importance |
|-----------------------|--------------|----------------------------------|-----------------------------|----------------------------|---------------------------|----------------------|-------------------|-------------------|------------------------------|---|--------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LTG | VPA | Relative (95% CI) | Absolute | | |
| 1 (Coppo- la 2004) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | serious ² | none | 7/19 (36.8%) | 12/19 (63.2%) | RR 0.58 (0.3 to 1.15) | 265 fewer per 1000 (from 442 fewer to 95 more) | ⊕⊕OO LOW | CRITICAL |
| Seizure free | edom - 12 m | nonths | | | | | | | | | | |
| 1 (Coppo- la 2004) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 10/19 (52.6%) | 13/19 (68.4%) | RR 0.77 (0.46 to 1.3) | 157 fewer per 1000 (from 369 fewer to 205 more) | ⊕⊕OO LOW | CRITICAL |
| Freedom fro | om treatme | nt failure a | t 16 or 20 weeks* | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 43/146 (29.5%) | 85/146 (58.2%) | RR 0.51 (0.38 to 0.67) | 285 fewer per 1000 (from 192 fewer to 361 fewer) | ⊕⊕⊕ HIGH | CRITICAL |
| Freedom fro | om treatme | nt failure a | t 12 months* | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 31/146 (21.2%) | 64/146 (43.8%) | RR 0.48 (0.34 to 0.7) | 228 fewer per 1000 (from 132 fewer to 289 fewer) | ⊕⊕⊕⊕ HIGH | CRITICAL |
| Adverse eve | ents - any (| follow-up 1 | 12 months) | | | | | | | | | |
| 1 (Coppo- la 2004) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 6/19 (31.6%) | 2/19 (10.5%) | RR 3 (0.69 to 13.03) | 211 more per 1000 (from 33 fewer to 1000 more) | ⊕⊕OO LOW | CRITICAL |
| Adverse eve | ents - serio | us (follow- | up 16 or 20 weeks | s) | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 2/149 (1.3%) | 2/147 (1.4%) | RR 0.99 (0.14 to 6.91) | 0 fewer per 1000 (from 12 fewer to 80 more) | ⊕⊕OO LOW | CRITICAL |

| Quality asso | essment | | | | | | Number o | of patients | Effect | | Quality | Importance |
|-----------------------|-------------|----------------------------------|-----------------------------|----------------------------|---------------------------|----------------------|-------------------|-------------------|--------------------------------|--|------------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LTG | VPA | Relative (95% CI) | Absolute | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 2/149 (1.3%) | 2/147 (1.4%) | RR 0.99 (0.14 to 6.91) | 0 fewer per 1000 (from 12 fewer to 80 more) | ⊕⊕OO LOW | CRITICAL |
| Treatment of | cessation d | ue to adve | rse events (follow | -up 12 months) | | | | | | | | |
| 1 (Coppo- la 2004) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ⁴ | none | 0/19 (0%) | 0/19 (0%) | RD 0.00 (- 0.10 to 0.10) | 0 fewer per 1000 (from 10 fewer to 10 more) | ⊕OOO VERY LOW | CRITICAL |
| Intolerable a | adverse eff | ects (follov | v-up 16 or 20 weel | ks) | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 25/146 (17.1%) | 35/146 (24%) | RR 0.71 (0.45 to 1.13) | 70 fewer per 1000 (from 132 fewer to 31 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Intolerable a | adverse eff | ects (follov | v-up 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 29/146 (19.9%) | 48/146 (32.9%) | RR 0.6 (0.41 to 0.9) | 132 fewer per 1000 (from 33 fewer to 194 fewer) | ⊕⊕⊕O MODERATE | CRITICAL |
| Study drug | discontinu | ed – no rea | son reported (foll | ow-un 16 or 20 v | weeks) | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 1/146 (0.68%) | 0/146 (0%) | RR 3 (0.12 to 73.04) | 10 more per 1000 (from 10 fewer to 30 more) | ⊕⊕OO LOW | CRITICAL |
| Withdrawal | from study | (follow-up | 3 months) | | | | | | | | | |
| 1 (Coppo- la 2004) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 6/19 (31.6%) | 3/19 (15.8%) | RR 2 (0.58 to 6.85) | 158 more per 1000 (from 66 fewer to 924 more) | ⊕OOO VERY LOW | CRITICAL |

| Quality ass | essment | | | | | | Number o | of patients | Effect | | Quality | Importance |
|---|------------|----------------------------------|--------------------------------|----------------------------|---------------------------|----------------------|-------------------|-------------------|------------------------------|---|------------------|----------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LTG | VPA | Relative (95% CI) | Absolute | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | very serious ³ | none | 18/146 (12.3%) | 15/146 (10.3%) | RR 1.2 (0.63 to 2.29) | 21 more per 1000 (from 38 fewer to 133 more) | ⊕⊕OO LOW | CRITICAL |
| Withdrawal | from study | (12 month | ıs) | | | | | | | | | |
| 2 (Coppo- la 2004, Glauser 2013) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 24/165 (14.5%) | 24/165 (14.5%) | RR 1 (0.59 to 1.69) | 0 fewer per 1000 (from 60 fewer to 100 more) | ⊕000 VERY LOW | CRITICAL |
| Conners' pe | erformance | test score | > 0.60 [¥] (follow-up | 16 or 20 weeks) | | | | | | | | |
| 1 (Glauser 2010) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 25/104 (24%) | 52/106 (49.1%) | RR 0.49 (0.33 to 0.73) | 250 fewer per 1000 (from 132 fewer to 329 fewer) | ⊕⊕⊕⊕ HIGH | IM- PORTANT |
| Conners' pe | erformance | test score | > 0.60 [¥] (follow-up | 12 months) | | | | | | | | |
| 1 (Glauser 2013) | RCT | no serious risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 8/30 (26.7%) | 34/61 (55.7%) | RR 0.48 (0.25 to 0.9) | 290 fewer per 1000 (from 56 fewer to 418 fewer) | ⊕⊕⊕O MODERATE | IM- PORTANT |

^{*} Treatment failure defined as persistence of absence seizures, a generalised tonic-clonic seizure at any time, excessive drug-related systemic toxicity of at least 3.0 from baseline, dose-limiting toxicity after a single downward dose modification, and withdrawal initiated by the parent or physician.

Table 11: Clinical evidence profile. Comparison 3: levetiracetam versus placebo

| Quality ass | essment | | | | | | Number o | f patients | Effect | | | |
|-------------|---------|---------|---------------|--------------|-------------|----------------|----------|------------|----------|----------|---------|------------|
| Number | Design | Risk of | Inconsistency | Indirectness | Imprecision | Other | LEV | Placebo | Relative | Absolute | | |
| of studies | | bias | | | | considerations | | | (95% CI) | | Quality | Importance |

^{*}An index of 0.60 corresponds to a 60% probability that the child has clinically significant attention deficit disorder 1 Serious risk of bias in the evidence contributing to the outcomes as per RoB 2

 ² 95% Confidence interval crosses 1 MID (0.8 or 1.25)
 ³ 95% Confidence interval crosses both MIDs (0.8 and 1.25)

⁴ Absolute effect range crosses 2 MIDs (10 more per 1000 and 10 fewer per 1000)

| Quality ass | essment | | | | | | Number o | of patients | Effect | | | |
|---------------------|--------------|----------------------|-----------------------------|----------------------------|---------------------------|----------------------|------------------|-----------------|--------------------------------|---|------------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LEV | Placebo | Relative (95% CI) | Absolute | Quality | Importance |
| Responder | status (free | from clini | cal seizures on da | ays 13 and 14, an | nd no EEG seizu | res during standar | d EEG on d | lay 14) – ITT | Γ analysis | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ² | none | 9/38 (23.7%) | 1/21 (4.8%) | RR 4.97 (0.68 to 36.61) | 189 more per 1000 (from 15 fewer to 1000 more) | ⊕OOO VERY LOW | CRITICAL |
| Patients fre | e from clini | ical and EE | G seizures - days | 4-7 (follow-up 4 | -7 days) | | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ² | none | 4/38 (10.5%) | 0/21 (0%) | POR 5.14 (0.63 to 42.06) | 110 more per 1000 (from 10 fewer to 220 more) | ⊕OOO VERY LOW | CRITICAL |
| Patients fre | e from clini | ical and EE | G seizures - days | 11-14 (follow-up | 11-14 days) | | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | serious ³ | none | 7/38 (18.4%) | 0/21 (0%) | POR 5.65 (1.1 to 28.96) | 180 more per 1000 (from 50 more to 320 more) | ⊕⊕OO LOW | CRITICAL |
| 50% reduct | ion (vs bas | eline) in tot | tal duration of EE | G seizures on da | y 14 (follow-up | mean 14 days) | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ² | none | 12/38 (31.6%) | 3/21 (14.3%) | RR 2.21 (0.7 to 6.96) | 173 more per 1000 (from 43 fewer to 851 more) | ⊕OOO VERY LOW | CRITICAL |
| Adverse ev | ents - any (| follow-up n | nean 14 days) | | | | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ² | none | 7/38 (18.4%) | 3/21 (14.3%) | RR 1.29 (0.37 to 4.47) | 41 more per 1000 (from 90 fewer to 496 more) | ⊕OOO VERY LOW | CRITICAL |
| Adverse ev | ents – seric | ous (follow- | -up mean 14 days |) | | | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ⁴ | none | 0/38 | 0/21 | RD 0.00 (- 0.07 to 0.07) | 0 more per 1000 (from 70 fewer to 70 more) | ⊕000 VERY LOW | CRITICAL |
| Adverse ev | ents - thou | ght to be re | elated to treatmen | t (follow-up mea | n 14 days) | | | | | | | |
| 1 (Fattore 2011) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ² | none | 3/38 (7.9%) | 0/21 (0%) | POR 4.99 (0.45 to 55.33) | 80 more per 1000 (from 30 fewer to 190 more) | ⊕000 VERY LOW | CRITICAL |
| Adverse ev | ents - leadi | ng to disco | ntinuation (follow | -up mean 14 day | ys) | | | | | | | |
| 1 (Fattore | RCT | serious ¹ | no serious | no serious | very serious ² | none | 1/38 | 0/21 | POR 4.72 | 30 more per | ⊕000 | CRITICAL |

| Quality ass | essment | | | | | | Number o | of patients | Effect | | | |
|-------------------|---------|--------------|---------------|--------------|-------------|----------------------|----------|-------------|----------------------|--|----------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LEV | Placebo | Relative (95% CI) | Absolute | Quality | Importance |
| 2011) | | | inconsistency | indirectness | | | (2.6%) | (0%) | (0.08 to 283.20) | 1000 (from 60 fewer to 110 more) | VERY LOW | |

Table 12: Clinical evidence profile. Comparison 4: lamotrigine versus placebo

| 10.010 12 | · Cimioui | - Tradition | promor comp | 2113011 4. IUIII0 | angino volo | ao piaceso | | | | | | |
|-------------------|--------------|----------------------|-----------------------------|-------------------------|---------------------------|----------------------|-----------------|-----------------|-----------------------------|---|---------------------|------------|
| Quality ass | essment | | | | | | Number | of patients | Effect | | | |
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | LTG | Placebo | Relative (95% CI) | Absolute | Quality | Importance |
| Remained s | seizure free | at end of pla | cebo controlled p | hase (follow-up | mean 4 weeks) | | | | | | | |
| 1 (Frank 1999) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | serious ² | none | 9/14 (64.3%) | 3/14 (21.4%) | RR 3 (1.02 to 8.8) | 429 more per 1000 (from 4 more to 1000 more) | ⊕⊕OO LOW | CRITICAL |
| Adverse ev | ents - leadi | ng to withdra | awal from study | | | | | | | | | |
| 1 (Frank 1999) | RCT | serious ¹ | no serious inconsistency | no serious indirectness | very serious ³ | none | 0/14 (0%) | 0/14 (0%) | RD 0.00 (- 0.13 to 0.13) | 0 more per 1000 (from 130 fewer to 130 more) | ⊕OOO VERY LOW | CRITICAL |

¹ Serious risk of bias in the evidence contributing to the outcomes as per RoB 2 ² Confidence interval crosses one MID (0.8 or 1.25)

Table 13: Clinical evidence profile. Comparison 5: ethosuximide versus lamotriqine

| Quality ass | essment | | | | | | Number o | of patients | Effect | | | |
|-------------------|------------|---------------|-------------------|--------------|-------------|----------------------|----------|-------------|----------------------|----------|---------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ETH | LTG | Relative (95% CI) | Absolute | Quality | Importance |
| Freedom fre | om treatme | nt failure at | t 16 or 20 weeks* | | | | | | | | | |

Serious risk of bias in the evidence contributing to the outcomes as per RoB 2
 Confidence intervals cross both MIDs (0.8 and 1.25)
 Confidence interval crosses one MID (0.8 or 1.25)
 Absolute effect range crosses 2 MIDs (10 more per 1000 and 10 fewer per 1000)

³ Absolute effect range crosses 2 MIDs (10 more per 1000 and 10 fewer per 1000)

| Quality ass | essment | | | | | | Number o | f patients | Effect | | | |
|---------------------|--------------|---------------------------------|-----------------------------|----------------------------|---------------------------|----------------------|-------------------|-------------------|-------------------------------|--|------------------|------------|
| Number of studies | Design | Risk of bias | Inconsistency | Indirectness | Imprecision | Other considerations | ETH | LTG | Relative (95% CI) | Absolute | Quality | Importance |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 81/154 (52.6%) | 43/146 (29.5%) | RR 1.79 (1.33 to 2.39) | 233 more per 1000 (from 97 more to 409 more) | ⊕⊕⊕⊕ HIGH | CRITICAL |
| Freedom fre | om treatme | nt failure a | t 12 months* | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | no serious imprecision | none | 70/154 (45.5%) | 31/146 (21.2%) | RR 2.14 (1.5 to 3.06) | 242 more per 1000 (from 106 more to 437 more) | ⊕⊕⊕⊕ HIGH | CRITICAL |
| Adverse ev | ents - serio | us (follow- | up 16 or 20 weeks |) | | | | | | | | |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | very serious ¹ | none | 4/155 (2.6%) | 2/149 (1.3%) | RR 1.92 (0.36 to 10.34) | 12 more per 1000 (from 9 fewer to 125 more) | ⊕⊕OO LOW | CRITICAL |
| Adverse ev | ents - serio | us (follow- | up 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | very serious ¹ | none | 4/155 (2.6%) | 2/149 (1.3%) | RR 1.92 (0.36 to 10.34) | 12 more per 1000 (from 9 fewer to 125 more) | ⊕⊕OO LOW | CRITICAL |
| Intolerable | adverse eff | ects (follov | v-up 16 or 20 weel | (s) | | | | | | | | |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 37/154 (24%) | 25/146 (17.1%) | RR 1.4 (0.89 to 2.21) | 68 more per 1000 (from 19 fewer to 207 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Intolerable | adverse eff | ects (follov | v-up 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 38/154 (24.7%) | 29/146 (19.9%) | RR 1.24 (0.81 to 1.9) | 48 more per 1000 (from 38 fewer to 179 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Study drug | discontinu | ed – no rea | son reporte (follo | w-up 16 or 20 we | eks) | | | | | | | |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | very serious ¹ | none | 0/154 (0%) | 1/146 (0.68%) | POR 0.13 (0.00 to 6.47) | 5 fewer per 1000 (from 7 fewer to 44 more) | ⊕⊕OO LOW | CRITICAL |

| Quality assessment Number Design Risk of Inconsistency Indirectness Imprecision Other | | | Number of patients | | Effect Relative Absolute | | | | | | | |
|--|-------------|---------------------------------|--------------------------|-------------------------------|---------------------------|----------------|-------------------|-------------------|------------------------------|--|------------------|----------------|
| of studies | Design | bias | moonsistemey | mancothess | Imprediction | considerations | | 2.0 | (95% CI) | Absolute | Quality | Importance |
| Withdrawal from study (follow-up 16 or 20 weeks) | | | | | | | | | | | | |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | very serious ¹ | none | 20/154 (13%) | 18/146 (12.3%) | RR 1.05 (0.58 to 1.91) | 6 more per 1000 (from 52 fewer to 112 more) | ⊕⊕OO LOW | CRITICAL |
| Withdrawal | from study | (follow-up | 12 months) | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 29/154 (18.8%) | 18/146 (12.3%) | RR 1.53 (0.89 to 2.63) | 65 more per 1000 (from 14 fewer to 201 more) | ⊕⊕⊕O MODERATE | CRITICAL |
| Conners' C | ontinuous I | Performano | e Test score > 0.6 | 60 [*] (follow-up 16 | or 20 weeks) | | | | | | | |
| 1 (Glauser 2010) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | serious ² | none | 35/106 (33%) | 25/104 (24%) | RR 1.37 (0.89 to 2.12) | 89 more per 1000 (from 26 fewer to 269 more) | ⊕⊕⊕O MODERATE | IM- PORTANT |
| Conners' Continuous Performance Test score > 0.60* (follow-up 12 months) | | | | | | | | | | | | |
| 1 (Glauser 2013) | RCT | no seri- ous risk of bias | no serious inconsistency | no serious indirectness | very serious ¹ | none | 20/70 (28.6%) | 8/30 (26.7%) | RR 1.07 (0.53 to 2.16) | 19 more per 1000 (from 125 fewer to 309 more) | ⊕⊕OO LOW | CRITICAL |

^{*}Treatment failure defined as persistence of absence seizures, a generalised tonic–clonic seizure at any time, excessive drug-related systemic toxicity of at least 3.0 from baseline, dose-limiting toxicity after a single downward dose modification, and withdrawal initiated by the parent or physician

*An index of 0.60 corresponds to a 60% probability that the child has clinically significant attention deficit disorder

¹ Confidence intervals cross both MIDs (0.8 and 1.25)

² MID crosses 1 MID (0.8 or 1.25)

1 Appendix G – Economic evidence study selection

- 2 Economic evidence study selection for review question: What antiseizure
- 3 therapies (monotherapy or add-on) are effective in the treatment of absence
- 4 seizures?
- 5 A single economic search was undertaken for all topics included in the scope of this guide-
- 6 line. See supplementary material 2 for further information.

1 Appendix H – Economic evidence tables

- 2 Economic evidence tables for review question: What antiseizure therapies (monotherapy or add-on) are effective in the
- 3 treatment of absence seizures?

5

4 No evidence was identified which was applicable to this review question.

1 Appendix I – Economic evidence profiles

- 2 Economic evidence profiles for review question: What antiseizure therapies (monotherapy or add-on) are effective in the
- 3 treatment of absence seizures?
- 4 No evidence was identified which was applicable to this review question.

1 Appendix J - Economic analysis

- 2 Economic evidence analysis for review question: What antiseizure therapies
- 3 (monotherapy or add-on) are effective in the treatment of absence seizures?
- 4 No economic analysis was conducted for this review question.

5

1 Appendix K - Excluded studies

- 2 Excluded clinical and economic studies for review question: What antiseizure
- 3 therapies (monotherapy or add-on) are effective in the treatment of absence
- 4 seizures?
- 5 Clinical studies

6 Table 14: Excluded studies and reasons for their exclusion

| Study | Reason for Exclusion |
|--|---|
| | |
| Actrn,, EpiNet-First Trial 2: comparison of efficacy of levetiracetam and sodium valproate in people with previously untreated epilepsy who have generalised seizures, Http://www.who.int/trialsearch/trial2.aspx? Trialid=actrn12615000556549, 2015 | Trial protocol |
| Actrn, EpiNet-First Trial 5: comparison of efficacy of levetiracetam and lamotrigine in people with previously untreated epilepsy who have unclassified seizures, and for whom sodium valporate is not deemed an acceptable anti-epileptic drug, Http://www.who.int/trialsearch/trial2.aspx? Trialid=actrn12615000641594, 2015 | Trial protocol |
| Actrn,, EpiNet-First Trial 4: comparison of efficacy of levetiracetam, lamotrigine and sodium valproate in people with previously untreated epilepsy who have unclassified seizures, Http://www.who.int/trialsearch/trial2.aspx? Trial-id=actrn12615000640505, 2015 | Trial protocol |
| Actrn, EpiNet-First Trial 3: comparison of efficacy of levetiracetam and lamotrigine in people with previously untreated epilepsy who have generalised seizures, and for whom sodium valporate is not deemed an acceptable anti-epileptic drug, Http://www.who.int/trialsearch/trial2.aspx? Trial-id=actrn12615000639527, 2015 | Trial protocol |
| Arya, R., Anand, V., Garg, S. K., Michael, B. D., Clobazam monotherapy for partial-onset or general- ized-onset seizures, Cochrane Database of Systemat- ic Reviews, 2014 (10) (no pagination), 2014 | Systematic review that focuses on people with new-onset focal or generalised seizures and does not report data on people with absence seizures or outcome data relating to this type of seizure |
| Arya, R., Giridharan, N., Anand, V., Garg, S. K., Clobazam monotherapy for focal or generalized sei- zures, Cochrane Database of Systematic Reviews, 2018 | Systematic review that focuses on people with new-onset focal or generalised seizures and does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Arzimanoglou, A., Rahbani, A., Zonisamide for the treatment of epilepsy, Expert Review of Neurotherapeutics, 6, 1283-1292, 2006 | Drug profile paper |
| Auvin, S., Advancing pharmacologic treatment options for pharmacologic treatment options for children with epilepsy, Expert Opinion on Pharmacotherapy, 17, 1475-1482, 2016 | Narrative review (no methodology reported) that does not report outcome data in sufficient detail for extraction extracted. The studies which are summarised have been checked for inclusion in this review |
| Basu, S., Bhattacharyya, K. B., Das, K., Das, D., Comparative study of sodium valproate and lamotrigi- | Conference abstract |

| Study | Reason for Exclusion |
|--|--|
| ne as monotherapy in the management of typical ab- | The state of the s |
| sence seizures, Epilepsia, 46, 277, Abstract no: p853, 2005 | |
| Benbadis, S., Klein, P., Schiemann, J., Diaz, A., Elmoufti, S., Whitesides, J., Efficacy, safety, and tolerability of brivaracetam with concomitant lamotrigine or concomitant topiramate in pooled Phase III randomized, double-blind trials: A post-hoc analysis, Epilepsy & Behavior, 80, 129-134, 2018 | Pooled analysis from RCTs reporting on interventions that do not meet the criteria specified in the protocol for this review |
| Beran, R. G., Berkovic, S. F., Dunagan, F. M., Vajda, F. J., Danta, G., Black, A. B., Mackenzie, R., Doubleblind, placebo-controlled, crossover study of lamotrigine in treatment-resistant generalised epilepsy, Epilepsia, 39, 1329-1333, 1998 | Mixed epilepsy population without sub- group analysis for people with absence seizures and data are not reported com- paratively |
| Besag, F. M. C., Wallace, S. J., Dulac, O., Alving, J., Spencer, S. C., Hosking, G., Lamotrigine for the treatment of epilepsy in childhood, Journal of Pediatrics, 127, 991-997, 1995 | Not comparative |
| Beydoun, A., D'Souza, J., Treatment of idiopathic generalized epilepsy - A review of the evidence, Expert Opinion on Pharmacotherapy, 13, 1283-1298, 2012 | Narrative review (no methodology reported) that does not report outcome data in sufficient detail for extraction extracted. The studies which are summarised have been checked for inclusion in this review |
| Biton, V., Preliminary open-label experience with to- piramate in primary generalized seizures, Epilepsia, 38, S42-S44, 1997 | Not comparative |
| Biton, V., Di Memmo, J., Shukla, R., Lee, Y. Y., Poverennova, I., Demchenko, V., Saiers, J., Adams, B., Hammer, A., Vuong, A., et al.,, Adjunctive lamotrigine XR for primary generalized tonic-clonic seizures in a randomized, placebo-controlled study, Epilepsy & Behavior, 19, 352― 358, 2010 | Mixed epilespy population without sub- group analysis for people with absence seizures |
| Biton, V., Shneker, B. F., Naritoku, D., Hammer, A. E., Vuong, A., Caldwell, P. T., Messenheimer, J. A., Longterm tolerability and safety of lamotrigine extended release: Pooled analysis of three clinical trials, Clinical Drug Investigation, 33, 359-364, 2013 | Mixed population without subgroup analysis for people with absence seizures |
| Bonnett, L. J., Tudur Smith, C., Smith, D., Williamson, P. R., Chadwick, D., Marson, A. G., Time to 12-month remission and treatment failure for generalised and unclassified epilepsy, Journal of neurology, neurosurgery, and psychiatry, 85, 603-610, 2014 | Describes development of a prognostic model |
| Brigo, F., Igwe, S. C., Ethosuximide, sodium valproate or lamotrigine for absence seizures in children and adolescents, Cochrane Database of Systematic Reviews, 2017 (2) (no pagination), 2017 | Systematic review. Included studies were checked for this review |
| Brigo, F., Igwe, S. C., Bragazzi, N. L., Lattanzi, S., Clonazepam monotherapy for treating people with newly diagnosed epilepsy, Cochrane Database of Sys- tematic Reviews, 2019 | Systematic review. Studies/data included in this paper had already been included in the evidence review, those not included were conference abstracts |
| Brigo, F., Igwe, S. C., Lattanzi, S., Ethosuximide, so- dium valproate or lamotrigine for absence seizures in children and adolescents, Cochrane Database of Sys- tematic Reviews, 2019 | Systematic review that has been updated – see other studies by same author |
| Buchanan, N., Lamotrigine in the treatment of absence seizures, Acta Neurologica Scandinavica, 92, 348, | Letter to the editor |

| Study | Reason for Exclusion |
|--|---|
| 1995 | |
| Bulau, P., Froscher, W., Schuchardt, V., Kreiten, K., A prospective randomised trial of the effectiveness of clonazepam and diazepam in petit mal epilepsy, Der nervenarzt, 57, 667― 671, 1986 | Not in English language |
| Bülau, P., Fröscher, W., Schuchardt, V., Kreiten, K., Prospective randomized study of the effectiveness of clonazepam and diazepam in petit mal status, Der nervenarzt, 57, 667― 671, 1986 | Not in English language |
| Buoni, S., Grosso, S., Fois, A., Lamotrigine treatment in childhood drug resistant epilepsy, Journal of Child Neurology, 13, 163-7, 1998 | Not comparative |
| Campos, M. S. A., Ayres, L. R., Morelo, M. R. S., Carizio, F. A. M., Pereira, L. R. L., Comparative efficacy of antiepileptic drugs for patients with generalized epileptic seizures: systematic review and network meta-analyses, International Journal of Clinical Pharmacy, 40, 589-598, 2018 | Studies included in this paper (relevant to absence seizures) had already been included in the evidence review |
| Cao, J., Lin, X. X., Ma, X. M., Liu, H., The efficacy and safety of lamotrigine for absence seizures in children and adolescents: A systematic review and meta-analysis, Journal of Clinical Neuroscience., 2019 | Systematic review that does not report outcome data in sufficient detail for extraction. The studies which are summarised have been checked for inclusion in this review |
| Chadwick, D., Does withdrawal of different antiepileptic drugs have different effects on seizure recurrence? Further results from the MRC Antiepileptic Drug Withdrawal Study, Brain, 122, 441-8, 1999 | Mixed population without subgroup analysis for people with absence seizures |
| Chandra, B., First seizure in adults: to treat or not to treat, Clinical Neurology & Neurosurgery, 94 Suppl, S61-3, 1992 | Mixed population without subgroup analysis for people with absence seizures |
| Cretin, B., Hirsch, E., Adjunctive antiepileptic drugs in adult epilepsy: how the first add-on could be the last, Expert Opinion on Pharmacotherapy, 11, 1053-67, 2010 | Narrative review (no methodology reported) that does not report outcome data in sufficient detail for extraction extracted. The studies which are summarised have been checked for inclusion in this review |
| Cross, J. H., Epilepsy (generalised seizures), BMJ clinical evidence, 2015 | Studies/data included in this paper (relevant to absence seizures) had already been included in the evidence review |
| Curatolo, P., Moavero, R., Lo Castro, A., Cerminara, C., Pharmacotherapy of idiopathic generalized epilepsies, Expert Opinion on Pharmacotherapy, 10, 5-17, 2009 | Narrative review (no methodology reported) that does not report outcome data. The studies which are summarised have been checked for inclusion in this review |
| Duchowny, M., Gilman, J., Messenheimer, J., et al., Long-term tolerability and efficacy of lamotrigine in pediatic patients with epilepsy, Journal of Child Neu- rology, 17, 278-285, 2002 | Not comparative |
| Euctr, F. R., Study to investigate the palatability, acceptability, pharmacokinetics, safety and tolerability, and treatment compliance of multidoses of ADV6770 as monotherapy or in combination, in children with childhood absence epilepsy, Http://www.who.int/trialsearch/trial2.aspx? Trialid=euctr2016-002313-22-fr, 2016 | Clinical trials registry entry |
| Fang, Y., Wu, X., Xu, L., Tang, X., Wang, J., Zhu, G., Hong, Z., Randomized-controlled trials of levetirace-tam as an adjunctive therapy in epilepsy of multiple | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of |

| Study | Reason for Exclusion |
|--|--|
| seizure types, Journal of Clinical Neuroscience, 21, | seizure |
| 55-62, 2014 | |
| French, J. A., Kanner, A. M., Bautista, J., et al, Efficacy and tolerability of the new antiepileptic drugs I: Treatment of new onset epilepsy. Report of the Therapeutics and Technology Assessment Subcommittee and Quality Standards Subcommittee of the American Academy of Neurology and the American Epilepsy Society, Neurology, 62, 1252-1260, 2004 | Systematic review that does not report outcome data in sufficient detail for extraction. The studies which are summarised have been checked for inclusion in this review |
| French, J. A., Kanner, A. M., Bautista, J., et al, Efficacy and Tolerability of the New Antiepileptic Drugs, I: Treatment of New-Onset Epilepsy: Report of the TTA and QSS Subcommittees of the American Academy of Neurology and the American Epilepsy Society, Epilepsia, 45, 401-409, 2004 | Systematic review that does not report outcome data in sufficient detail for extraction. The studies which are summarised have been checked for inclusion in this review |
| French, J. A., Kanner, A. M., Bautista, J., et al, Appendix C: Efficacy and tolerability of the new antiepileptic drugs I: Treatment of new onset epilepsy: Report of the Therapeutics and Technology Assessment Subcommittee and Quality Standards Subcommittee of the American Academy of Neurology and the American Epilepsy Society, CONTINUUM Lifelong Learning in Neurology, 13, 203-211, 2007 | Systematic review that does not report outcome data in sufficient detail for extraction. The studies which are summarised have been checked for inclusion in this review |
| Gamble, C., Williamson, P. R., Chadwick, D. W., Marson, A. G., A meta-analysis of individual patient responses to lamotrigine or carbamazepine monotherapy, Neurology, 66, 1310-1317, 2006 | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Gasparini, S., Ferlazzo, E., Giussani, G., Italiano, D., Cianci, V., Sueri, C., Spina, E., Beghi, E., Aguglia, U., Rapid versus slow withdrawal of antiepileptic monotherapy in 2-year seizure-free adult patients with epilepsy (RASLOW) study: a pragmatic multicentre, prospective, randomized, controlled study, Neurological Sciences, 37, 579-583, 2016 | Trial protocol |
| Giorgi, L., Gomez, G., O'Neill, F., Hammer, A. E., Risner, M., The tolerability of lamotrigine in elderly patients with epilepsy, Drugs & Aging, 18, 621-30, 2001 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| Glauser, T., Ben-Menachem, E., Bourgeois, B., Cnaan, A., Guerreiro, C., Kalviainen, R., Mattson, R., French, J. A., Perucca, E., Tomson, T., Updated ILAE evidence review of antiepileptic drug efficacy and ef- fectiveness as initial monotherapy for epileptic sei- zures and syndromes, Epilepsia, 54, 551-563, 2013 | Systematic review that does not report outcome data in sufficient detail for extraction. The studies which are summarised have been checked for inclusion in this review |
| Hemery, C., Ryvlin, P., Rheims, S., Prevention of generalized tonic-clonic seizures in refractory focal epilepsy: A meta-analysis, Epilepsia, 55, 1789-1799, 2014 | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Hong, Z., Inoue, Y., Liao, W., Meng, H., Wang, X., Wang, W., Zhou, L., Zhang, L., Du, X., Tennigkeit, F., Efficacy and safety of adjunctive lacosamide for the treatment of partial-onset seizures in Chinese and Japanese adults: a randomized, double-blind, place-bo-controlled study, Epilepsy research, 127, 267― 275, 2016 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| Houtkooper, M. A., Lammertsma, A., Meyer, J. W., Goedhart, D. M., Meinardi, H., van Oorschot, C. A., Blom, G. F., Hoppener, R. J., Hulsman, J. A., Ox- | Interventions do not meet the criteria specified in the protocol for this review |

| Study | Reason for Exclusion |
|---|---|
| carbazepine (GP 47.680): a possible alternative to | |
| carbamazepine?, Epilepsia, 28, 693-8, 1987 | |
| Huang, T. S., Zhu, J. L., Li, B., Hu, Y., Chen, L., Liao, J. X., Valproic acid versus lamotrigine as a monotherapy for absence epilepsy in children, Zhongguo dang dai er ke za zhi [Chinese journal of contemporary pediatrics], 11, 653― 655, 2009 | Not in English language |
| Kaminow, L., Schimschock, J. R., Hammer, A. E., Vuong, A., Lamotrigine monotherapy compared with carbamazepine, phenytoin, or valproate monotherapy in patients with epilepsy, Epilepsy & Behavior, 4, 659-66, 2003 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| Kasteleijn-Nolst Trenite, D., Genton, P., Brandt, C., Reed, R. C., The 'Photosensitivity Model' is (also) a model for focal (partial) seizures, Epilepsy Research, 133, 113-120, 2017 | Not a treatment study or systematic review of treatment |
| Kerr, M. P., Baker, G. A., Brodie, M. J., A randomized, double-blind, placebo-controlled trial of topiramate in adults with epilepsy and intellectual disability: Impact on seizures, severity, and quality of life, Epilepsy and Behavior, 7, 472-480, 2005 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| Krauss, G. L., Efficacy and tolerability of the new antiepileptic drugs I: treatment of new onset epilepsy: report of the Therapeutics and Technology Assessment Subcommittee and Quality Standards Subcommittee of the American Academy of Neurology and the American Epilepsy Society, Neurology, 64, 172-4; author reply 172-4, 2005 | Commentary |
| Kumar, S., Ramanujam, B., Chandra, P. S., Dash, D., Mehta, S., Anubha, S., Appukutan, R., Rana, M. K., Tripathi, M., Randomized controlled study comparing the efficacy of rapid and slow withdrawal of antiepileptic drugs during long-term video-EEG monitoring, Epilepsia, 59, 460-467, 2018 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| Marson, A. G., Maguire, M., Ramaratnam, S., Epilepsy, BMJ clinical evidence, 2009 | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Marson, A., Jacoby, A., Johnson, A., Kim, L., Gamble, C., Chadwick, D., Immediate versus deferred antiepileptic drug treatment for early epilepsy and single seizures: A randomised controlled trial, Lancet, 365, 2007-2013, 2005 | Systematic review that does not meet the inclusion criteria (evaluates immediate versus deferred treatment and does not report subgroup analysis for people with absence seizures). Included studies were checked for this review |
| Mattson, R. H., Cramer, J. A., Collins, J. F., McCutcheon, C. B., Fish, S. L., Mamdani, M. B., Rubino, F. A., Davenport, J., Lubozynski, M. F., Ramsay, R. E., Carter, G. S., Rowan, A. J., Browne, T. R., Ebersole, J. S., Treiman, D. M., Warner, J. J., Wilder, B. J., Salinsky, M., Arroyo, Y., A comparison of valproate with carbamazepine for the treatment of complex partial seizures and secondarily generalized tonic-clonic seizures in adults, New England Journal of Medicine, 327, 765-771, 1992 | Mixed epilepsy population without sub- group analysis for people with absence seizures |
| McAuley, C., McShane, T., Ethosuximide was superior to valproate and lamotrigine in controlling absence seizures and minimising side effects, Archives of Dis- | Conference abstract |

| Study | Reason for Exclusion |
|--|---|
| ease in Childhood: Education and Practice Edition, 96, 119, 2011 | |
| McDonald, T. J. W., Henry-Barron, B. J., Felton, E. A., Gutierrez, E. G., Barnett, J., Fisher, R., Lwin, M., Jan, A., Vizthum, D., Kossoff, E. H., Cervenka, M. C., Improving compliance in adults with epilepsy on a modified Atkins diet: A randomized trial, Seizure, 60, 132-138, 2018 | Interventions do not meet the criteria specified in the protocol for this review |
| Messenheimer, J.A., Giorgi, L., Risner, M.E., The tolerability of lamotrigine in children, Drug Safety, 22, 303-312, 2000 | Systematic review. Studies/data included in this paper had already been included in the evidence review, data not included were taken from a clinical database and insufficient detail is provided to allow extraction of results |
| Miura, H., Shirai, H., Sunaoshi, W., Effectiveness and plasma levels of clonazepam in the treatment of absence seizures, Journal of the japan epilepsy society, 5, 41― 49, 1987 | Not in English language |
| Moon, K. T., What is the best treatment for childhood absence epilepsy?, American Family Physician, 83, 81-82, 2011 | Not primary research or a systematic review |
| Nadkarni, S., LaJoie, J., Devinsky, O., Current treatments of epilepsy, Neurology, 64, S2-S11, 2005 | Narrative review, does not summarise studies in which people with absence seizures were included |
| Nct., IV Keppra in the Emergency Department for Prevention of Early Recurrent Seizures, https://clinicaltrials.gov/show/NCT00510783, 2007 | Trial registry entry |
| Nct., Does Gabapentin and Lamotriginel Have Significantly Fewer Side-Effects While Providing Equal or Better Seizure Control Than the Current Drug Choice, Carbamazepine, for the Treatment of Seizures in the Elderly, https://clinicaltrials.gov/show/NCT00007670, 2000 | Trial registry entry |
| Nct,, Modified Atkins Diet Plus KetoCal for Adult Epilepsy, https://clinicaltrials.gov/show/NCT01834482, 2013 | Trial registry entry |
| Nevitt, S. J., Sudell, M., Weston, J., Tudur Smith, C., Marson, A. G., Antiepileptic drug monotherapy for epilepsy: a network meta-analysis of individual participant data, Cochrane Database of Systematic Reviews, 2017 | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Nevitt, S. J., Sudell, M., Weston, J., Tudur Smith, C., Marson, A. G., Antiepileptic drug monotherapy for epilepsy: A network meta-analysis of individual participant data, Cochrane Database of Systematic Reviews, 2017 (6) (no pagination), 2017 | Systematic review that does not report data on people with absence seizures/outcome data relating to this type of seizure |
| Nieto-Barrera, M., Characteristics and indications of topiramate, Revista de neurologia, 35, S88-S95, 2002 | Not in English language |
| Ohtahara, S., Zonisamide in the management of epilepsy - Japanese experience, Epilepsy Research, 68, S25-S33, 2006 | Systematic review which included non- comparative studies (outcomes relating to absence studies based on survey data) |
| Ormrod, D., McClellan, K., Topiramate: A review of its use in childhood epilepsy, Paediatric Drugs, 3, 293-319, 2001 | Mixed epilepsy population without sub- group analysis for people with absence seizures (other than in the context of Len- nox Gastaut syndrome) |
| Pohlmann-Eden, B., Marson, A. G., Noack-Rink, M., et | Mixed epilepsy population without sub- |

| Study | Reason for Exclusion |
|--|---|
| al., Comparative effectiveness of levetiracetam, valproate and carbamazepine among elderly patients with newly diagnosed epilepsy: subgroup analysis of the randomized, unblinded KOMET study, BMC Neurology, 16, 149, 2016 | group analysis for people with absence seizures |
| Posner, E., Absence seizures in children, Clinical Evidence, 307-13, 2004 | Unavailable from the British Library (checked 18/03/21) |
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| Study | Reason for Exclusion |
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3 Economic studies

No economic evidence was identified for this review. See supplementary material 2 for further information.

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1 Appendix L - Research recommendations

- 2 Research recommendations for review question: What antiseizure therapies
- 3 (monotherapy or add-on) are effective in the treatment of absence seizures?
- 4 No research recommendations were made for this review question.