Clinical trial results:

A Double-Blind, Randomized, Multicenter, Placebo-controlled, In-Patient, Maximum 34 Day Study of Levetiracetam Oral Solution (20-50 mg/kg/day) as Adjunctive Treatment of Refractory Partial Onset Seizures in Pediatric Epileptic Subjects Ranging in Age from 1 Month to Less Than 4 Years of Age

Summary

EudraCT number	2004-000199-14	
Trial protocol	CZ GB BE HU IT	
Global end of trial date	26 January 2007	
Results information		
Result version number	v1 (current)	
This version publication date	28 June 2016	
First version publication date	22 July 2015	

Trial information

Trial identification	
Sponsor protocol code	N01009
Additional study identifiers	
ISRCTN number	-
ClinicalTrials.gov id (NCT number)	NCT00175890
WHO universal trial number (UTN)	-
Notes:	

Sponsors

Sponsor organisation name	UCB, Inc.
Sponsor organisation address	1950 Lake Park Drive, Smyrna, United States, 30080
Public contact	Clinical Trial Registries and Results Disclosure, UCB BIOSCIENCES GmbH, +49 2173 48 15 15, clinicaltrials@ucb.com
Scientific contact	Clinical Trial Registries and Results Disclosure, UCB BIOSCIENCES GmbH, +49 2173 48 15 15, clinicaltrials@ucb.com

Notes:

Paediatric regulatory details

Is trial part of an agreed paediatric investigation plan (PIP)	No
Does article 45 of REGULATION (EC) No 1901/2006 apply to this trial?	No
Does article 46 of REGULATION (EC) No 1901/2006 apply to this trial?	Yes
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Notes:

Results analysis stage	
Analysis stage	Final
Date of interim/final analysis	22 March 2007
Is this the analysis of the primary completion data?	Νο
Global end of trial reached?	Yes
Global end of trial date	26 January 2007
Was the trial ended prematurely?	Νο
Notes:	

General information about the trial

Main objective of the trial:

To evaluate the efficacy and safety of levetiracetam (LEV) used as adjunctive treatment in pediatric subjects age 1 month to less than 4 years with refractory partial onset seizures.

Protection of trial subjects:

Not applicable	
Background therapy:	
Not applicable	
Evidence for comparator:	
Not applicable	
Actual start date of recruitment	15 October 2004
Long term follow-up planned	Yes
Long term follow-up rationale	Ethical reason
Long term follow-up duration	30 Months
Independent data monitoring committee (IDMC) involvement?	No

Notes:

Population of trial subjects

Subjects enrolled per country

Belgium: 2
Brazil: 26
Czech Republic: 7
France: 6
Germany: 15
Hungary: 3
Italy: 5
Mexico: 6
Poland: 1
Romania: 4
Russian Federation: 7
United Kingdom: 1
United States: 33
116
44

Subjects enrolled per age group	
In utero	0
Preterm newborn - gestational age < 37 wk	0
Newborns (0-27 days)	0
Infants and toddlers (28 days-23 months)	61
Children (2-11 years)	55
Adolescents (12-17 years)	0
Adults (18-64 years)	0
From 65 to 84 years	0
85 years and over	0

Subject disposition

Recruitment

Recruitment details:

This double-blind, randomized, multicenter, placebo-controlled, in-Patient study started recruiting in October 2004.

Pre-assignment

Screening details:

The Intent-to-treat (ITT) Population consisted of all randomized subjects who took at least one dose of study drug.

Period 1

Period 1 title	Overall Study (overall period)
Is this the baseline period?	Yes
Allocation method	Randomised - controlled
Blinding used	Double blind
Roles blinded	Subject, Investigator

Arms

Are arms mutually exclusive?	Yes
Arm title	Placebo

Arm description:

Matching oral solution to Levetiracetam b.i.d. (twice a day) for a maximum treatment duration of 20 days.

Arm type	Placebo
Investigational medicinal product name	Placebo
Investigational medicinal product code	РВО
Other name	
Pharmaceutical forms	Oral solution
Routes of administration	Oral use

Dosage and administration details:

Placebo solution, which is indistinguishable from the Levetiracetam oral solution.

Arm title	Levetiracetam

Arm description:

10 % oral solution Levetiracetam b.i.d. (twice a day) for a maximum treatment duration of 20 days.

Arm type	Experimental
Investigational medicinal product name	Levetiracetam (LEV)
Investigational medicinal product code	LEV
Other name	Keppra
Pharmaceutical forms	Oral solution
Routes of administration	Oral use

Dosage and administration details:

Dosing was stratified by age. A dose of 20 mg/kg/day titrating to 40 mg/kg/day for children one month to less than six months old and a dose of 25 mg/kg/day titrating to 50 mg/kg/day for children 6 month to less than 4 years old, was used in this study. The total daily dose was administered b.i.d.

Number of subjects in period 1	Placebo	Levetiracetam
Started	56	60
Completed	53	58
Not completed	3	2
Consent withdrawn by subject	1	-
AE, non-serious non-fatal	-	2
SAE, non-fatal	1	-
Protocol deviation	1	-

Baseline characteristics

Reporting groups		
Reporting group title	Placebo	
Reporting group description:		
Matching oral solution to Levetiracetam b.i.d. (twice a day) for a maximum treatment duration of 20 days.		
Reporting group title	Levetiracetam	
Reporting group description:		
10 % oral solution Levetiracetam b.i.d. (twice a day) for a maximum treatment duration of 20 days.		

Placebo Levetiracetam Total **Reporting group values** Number of subjects 60 116 56 Age categorical Units: Subjects 0 - < = 27 days0 0 0 28 days - < 24 months 29 32 61 24 months - < 12 years 27 28 55 Age Continuous Units: months arithmetic mean 23.46 23.4 standard deviation ± 12.06 ± 13.43 Gender Categorical Units: Subjects Male 27 30 57 Female 29 30 59 Race/Ethnicity, Customized Units: Subjects Caucasian 39 54 93 American Indian or Alaska Native 2 4 6 2 10 Other/mixed race 8 Black 6 0 6 Asian 1 0 1 Race/Ethnicity, Customized Units: Subjects Hispanic or Latino 16 22 38 Not Hispanic or Latino 78 40 38

Primary: Responder Rate for total partial onset seizures as computed from the 48hour Evaluation video-EEG (post-baseline) and the 48-hour Selection video-EEG (baseline)

End point title	Responder Rate for total partial onset seizures as computed
	from the 48-hour Evaluation video-EEG (post-baseline) and the
	48-hour Selection video-EEG (baseline)

End point description:

Responder Rate is defined as the number of subjects with a 50 % reduction from baseline in their Average Daily Frequency (ADF) for partial onset seizures divided by the total number of subjects. If a subject had < 24 hours of usable Evaluation video-EEG time (including zero time available) and withdrawal from the study with reasons linked to lack or loss of efficacy, the subject was counted as a non-responder.

End point typePrimaryEnd point timeframe:

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	51	58	
Units: percentage of participants			
number (not applicable)			
Responder (percentage)	19.6	43.1	
Non-Responder (percentage)	80.4	56.9	

Statistical analyses

Statistical analysis title	Statistical Analysis 1
Comparison groups	Modified Intent-to-Treat (LEV treated Subjects) v Modified Intent-to-Treat (Placebo treated Subjects)
Number of subjects included in analysis	109
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.013
Method	Fisher exact
Parameter estimate	Odds ratio (OR)
Point estimate	3.11
Confidence interval	
level	95 %
sides	2-sided
lower limit	1.22
upper limit	8.26

Secondary: Responder rate for total seizures (all types) as computed from the 48-

hour Evaluation video-EEG (post-baseline) and the 48-hour Selection video-EEG (baseline)

End point title	Responder rate for total seizures (all types) as computed from the 48-hour Evaluation video-EEG (post-baseline) and the 48-
	hour Selection video-EEG (baseline)

End point description:

Responder Rate is defined as the number of subjects with a 50 % reduction from baseline in their Average Daily Frequency (ADF) for all seizure types divided by the total number of subjects. Subjects who withdrew or dropped out before the first 24 hours Evaluation video-EEG with reasons linked to lack of efficacy were considered as non-responders.

All (total) seizures were defined as the total of Type I (partial onset) + Type II (Primary generalized) + Type III (unclassified epileptic).

End point type

Secondary

End point timeframe:

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	51	58	
Units: percentage of participants			
number (not applicable)			
Responder (percentage)	19.6	43.1	
Non-Responder (percentage)	80.4	56.9	

Statistical analyses

No statistical analyses for this end point

Secondary: Percent reduction in Average Daily Frequency (ADF) of partial onset seizures recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG

End point title	Percent reduction in Average Daily Frequency (ADF) of partial onset seizures recorded on the 48-hour Evaluation video-EEG
	compared to those recorded on the 48-hour Selection video- EEG

End point description:

A positive value in Percent reduction from Selection Period to Evaluation Period indicates an improvement.

 End point type
 Secondary

 End point timeframe:
 Secondary

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	50	55	
Units: Percent reduction			
arithmetic mean (standard deviation)			
arithmetic mean (standard deviation)	-20.93 (± 111.47)	24.98 (± 91.49)	

Statistical analyses

No statistical analyses for this end point

Secondary: Percent reduction in Average Daily Frequency (ADF) of total seizures (all types) recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG

End point title	Percent reduction in Average Daily Frequency (ADF) of total seizures (all types) recorded on the 48-hour Evaluation video- EEG compared to those recorded on the 48-hour Selection video-EEG
End point description:	

End point description:

A positive value in Percent reduction from Selection Period to Evaluation Period indicates an improvement.

All (total) seizures were defined as the total of Type I (partial onset) + Type II (Primary generalized) + Type III (unclassified epileptic).

End point type	Secondary
End point timeframe:	

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	50	55	
Units: Percent reduction			
arithmetic mean (standard deviation)			
arithmetic mean (standard deviation)	-20.93 (± 111.47)	25.08 (± 91.56)	

Statistical analyses

No statistical analyses for this end point

Secondary: Absolute reduction in Average Daily Frequency (ADF) of partial onset

seizures recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG

End point title Absolute reduction in Average Daily Frequency (ADF) of partial onset seizures recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG

End point description:

A positive value in Absolute reduction from Selection Period to Evaluation Period indicates an improvement.

End point type

Secondary

End point timeframe:

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	51	58	
Units: Absolute reduction			
arithmetic mean (standard deviation)			
arithmetic mean (standard deviation)	-0.86 (± 13.81)	8.54 (± 25.67)	

Statistical analyses

No statistical analyses for this end point

Secondary: Absolute reduction in Average Daily Frequency (ADF) of total seizures (all types) recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG

End point title	Absolute reduction in Average Daily Frequency (ADF) of total seizures (all types) recorded on the 48-hour Evaluation video-
	EEG compared to those recorded on the 48-hour Selection video-EEG

End point description:

A positive value in Absolute reduction from Selection Period to Evaluation Period indicates an improvement.

All (total) seizures were defined as the total of Type I (partial onset) + Type II (Primary generalized) + Type III (unclassified epileptic).

End point type	Secondary
End point timeframe:	

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Modified Intent-to-Treat (Placebo treated Subjects)	Modified Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	51	58	
Units: Absolute reduction			
arithmetic mean (standard deviation)			
arithmetic mean (standard deviation)	-0.86 (± 13.81)	9.12 (± 26.35)	

Statistical analyses

No statistical analyses for this end point

Secondary: Percent reduction in Average Daily Frequency (ADF) of electro-clinical partial onset seizures recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48-hour Selection video-EEG for children 1 month to less than 6 months old

End point title	Percent reduction in Average Daily Frequency (ADF) of electro- clinical partial onset seizures recorded on the 48-hour Evaluation video-EEG compared to those recorded on the 48- hour Selection video-EEG for children 1 month to less than 6 months old
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End point description:

A positive value in Percent reduction from Selection Period to Evaluation Period indicates an improvement.

For children 1 month to less than 6 months old, partial onset seizure counts were based on electroclinical seizures plus electrographic seizures.

End point type

Secondary

End point timeframe:

48-hours in Evaluation Period and 48-hours in Selection Period

End point values	Subset of mITT (Placebo treated Subjects)	Subset of mITT (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	4	3	
Units: Percent reduction			
arithmetic mean (standard deviation)			
arithmetic mean (standard deviation)	28.46 (± 44.89)	63.72 (± 28.41)	

Statistical analyses

No statistical analyses for this end point

Secondary: Percentage of drop-outs for any reasons during the study

End point title	Percentage of drop-outs for any reasons during the study
End noint description.	

End point description:

End point type Secondary

End point timeframe:

During the study (up to 20 days)

End point values	Intent-to-Treat (Placebo treated Subjects)	Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	56	60	
Units: percentage of participants			
number (not applicable)			
Drop-outs (percentage)	5.4	3.3	

Statistical analyses

No statistical analyses for this end point

Secondary: Percentage of drop-outs due to lack of efficacy during the study

End point title	Percentage of drop-outs due to lack of efficacy during the study
End point description:	

End point type	Secondary
End point timeframe:	
During the study (up to 20 days)	

End point values	Intent-to-Treat (Placebo treated Subjects)	Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	56	60	
Units: percentage of participants			
number (not applicable)			
Drop-outs (percentage)	0	0	

Statistical analyses

Secondary: Percentage of drop-outs before 24 hours of Evaluation video-EEG for reasons other than lack or loss of efficacy

End point title

Percentage of drop-outs before 24 hours of Evaluation video-EEG for reasons other than lack or loss of efficacy

End point description:

End point type

Secondary

End point timeframe:

During the study (up to 20 days)

End point values	Intent-to-Treat (Placebo treated Subjects)	Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	56	60	
Units: percentage of participants			
number (not applicable)			
Drop-outs (percentage)	3.6	1.7	

Statistical analyses

No statistical analyses for this end point

Secondary: Time to Exit (TTE) during the Evaluation Period

End point title

Time to Exit (TTE) during the Evaluation Period

End point description:

For early termination subjects in the Evaluation period the TTE is the time to discontinuing the study for any reason. TTE was defined as the day of study discontinuation – the day of randomization + 1. For completed subjects, the TTE was censored on Day 6.

End point type	Secondary	
End point timeframe:		
During Evaluation Period (Day 1 to Day 6	5)	

End point values	Intent-to-Treat (Placebo treated Subjects)	Intent-to-Treat (LEV treated Subjects)	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	56 ^[1]	60 ^[2]	
Units: Days			
median (confidence interval 95%)			
Median (95 % CI)	9999 (999 to 99999)	9999 (999 to 99999)	

Notes:

- [1] 999 / 9999 / 99999 = statistic not estimable
- [2] 999 / 9999 / 99999 = statistic not estimable

Statistical analyses

No statistical analyses for this end point

Adverse events information

Timeframe for reporting adverse events:

Adverse Events were collected from Selection Period (Day -8 to Day 0) until Post Treatment Follow-up (Day 24 \pm 1).

Adverse event reporting additional description:

Adverse Events refer to the Intent-to-treat (ITT) Population, including all randomized subjects who took at least one dose of study drug.

Assessment type	Non-systematic		
Dictionary used			
Dictionary name	MedDRA		
Dictionary version	9.0		
Reporting groups			
Reporting group title	Levetiracetam		
Reporting group description:			
10 % oral solution Levetiracetam b.i.	d. (twice a day) for a maximum treatment duration of 20 days.		
Reporting group title	Placebo		
Reporting group description:			

Matching oral solution to Levetiracetam b.i.d. (twice a day) for a maximum treatment duration of 20 days.

Serious adverse events	Levetiracetam	Placebo	
Total subjects affected by serious adverse events			
subjects affected / exposed	1 / 60 (1.67%)	1 / 56 (1.79%)	
number of deaths (all causes)	0	0	
number of deaths resulting from adverse events	0	0	
General disorders and administration site conditions			
Pyrexia			
subjects affected / exposed	1 / 60 (1.67%)	0 / 56 (0.00%)	
occurrences causally related to treatment / all	0 / 1	0/0	
deaths causally related to treatment / all	0 / 0	0/0	
Infections and infestations			
Urinary tract infection			
subjects affected / exposed	0 / 60 (0.00%)	1 / 56 (1.79%)	
occurrences causally related to treatment / all	0/0	0 / 1	
deaths causally related to treatment / all	0/0	0/0	

Frequency threshold for reporting non-serious adverse events: 5 %

Levetiracetam	Placebo	
10 / 60 (21 67%)	14 / 56 (25 00%)	
19700(31.07%)	147 50 (25.00%)	
8 / 60 (13.33%)	1 / 56 (1.79%)	
8	1	
7 / 60 (11.67%)	0 / 56 (0.00%)	
7	0	
2 / 60 (3.33%)	4 / 56 (7.14%)	
2	4	
2 / 60 (3.33%)	3 / 56 (5.36%)	
2	4	
2/60(2.22%)	2 / 56 (5 26%)	
2 / 00 (3.33%)	3750(5.30%)	
3	3	
1 / 60 (1.67%)	3 / 56 (5.36%)	
2	3	
1 / 60 (1.67%)	3 / 56 (5.36%)	
1	3	
	Levetiracetam 19 / 60 (31.67%) 8 / 60 (13.33%) 8 7 / 60 (11.67%) 7 2 / 60 (3.33%) 2 2 / 60 (3.33%) 2 2 / 60 (3.33%) 3 1 / 60 (1.67%) 2 1 / 60 (1.67%) 1	Levetiracetam Placebo 19 / 60 (31.67%) 14 / 56 (25.00%) 8 / 60 (13.33%) 1 / 56 (1.79%) 8 1 7 / 60 (11.67%) 0 / 56 (0.00%) 7 0 2 / 60 (3.33%) 4 / 56 (7.14%) 2 / 60 (3.33%) 3 / 56 (5.36%) 2 / 60 (3.33%) 3 / 56 (5.36%) 2 / 60 (3.33%) 3 / 56 (5.36%) 3 3 1 / 60 (1.67%) 3 / 56 (5.36%) 1 / 60 (1.67%) 3 / 56 (5.36%) 1 / 60 (1.67%) 3 / 56 (5.36%) 3 3

Substantial protocol amendments (globally)

Date	Amendment
01 April 2004	Amendment 1 dated 01-Apr-2004 modified the study design to incorporate neuropsychological testing, lengthened the down-titration period to decrease the risk of withdrawal seizures, lengthened the enrollment period and clarified the dosing scheme.
14 February 2005	Amendment 2 dated 14-Feb-2005 defined the policy regarding the enrollment of infants Born pre-term (before 37 weeks gestational age), defined the minimum weight of 4.0 kg and clarified the requirement for pre-screening of subjects by the UCB Clinical Research Physician (CRP).
14 June 2006	Amendment 3 dated 14-Jun-2006 added Argentina, Hungary, Poland, Romania and Russia, updated the background information to reflect FDA and European Commission approval of Keppra® for partial onset seizures in children ages four and above, and studies N159 and N157, defined the distinctive difference between electro-clinical seizure and electrographic seizures, and provided clarity to the description of cluster seizures.

Were there any global substantial amendments to the protocol? Yes

Notes:

Interruptions (globally)

Were there any global interruptions to the trial? No

Limitations and caveats

None reported

Online references

http://www.ncbi.nlm.nih.gov/pubmed/19243423