

**HIGHLIGHTS OF PRESCRIBING INFORMATION**

These highlights do not include all the information needed to use levetiracetam safely and effectively. See full prescribing information for levetiracetam.

**Levetiracetam Tablets, USP** for oral use  
**Initial U.S. Approval: 1999**

----- **RECENT MAJOR CHANGES** -----  
 Indications and Usage, Partial Onset Seizures (1.1) [12/2011]  
 Dosage and Administration, Partial Onset Seizures (1.1) [12/2011]  
 Warnings and Precautions (6.1, 5.3, 5.4, 5.7, 5.8, 5.9) [12/2011]

----- **INDICATIONS AND USAGE** -----  
 Levetiracetam is an antiepileptic drug indicated for adjunctive therapy in the treatment of:  
 • Partial onset seizures in patients 4 years of age and older with epilepsy (1.1)  
 • Myoclonic seizures in patients 12 years of age and older with juvenile myoclonic epilepsy (1.2)  
 • Primary generalized tonic-clonic seizures in patients 6 years of age and older with idiopathic generalized epilepsy (1.3)

----- **DOSAGE AND ADMINISTRATION** -----  
 Use the oral solution for pediatric patients with body weight < 20 kg (4.4).  
 For pediatric patients, use weight-based dosing for the oral solution with a calibrated measuring device (not a household teaspoon or tablespoon) (2.1)

**Partial Onset Seizures**  
 • 4 Years To < 16 Years: 10 mg/kg twice daily, increase in increments of 10 mg/kg twice daily every 2 weeks to recommended dose of 50 mg/kg twice daily (2.2)  
 • Adults 16 Years And Older: 500 mg twice daily, increase as needed and tolerated in increments of 500 mg twice daily every 2 weeks to a maximum recommended dose of 1500 mg twice daily (2.2)

**Myoclonic Seizures in Adults and Pediatric Patients 12 Years and Older**  
 • 500 mg twice daily, increase by 500 mg twice daily every 2 weeks to recommended dose of 1500 mg twice daily (2.2)

**Primary Generalized Tonic-Clonic Seizures**  
 • 6 Years To < 16 Years: 10 mg/kg twice daily, increase in increments of 10 mg/kg twice daily every 2 weeks to recommended dose of 50 mg/kg twice daily (2.4)  
 • Adults 16 Years And Older: 500 mg twice daily, increase by 500 mg twice daily every 2 weeks to recommended dose of 1500 mg twice daily (2.4)

**Use With Other Antiepileptic Drugs**  
 • 500 mg twice daily, increase by 500 mg twice daily every 2 weeks to recommended dose of 1500 mg twice daily (2.4)

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**2.4 Primary Generalized Tonic-Clonic Seizures**

**Adults 16 Years And Older**  
 Treatment should be initiated with a dose of 1000 mg/day, given as twice-daily dosing (500 mg twice daily). Dosage should be increased by 1000 mg/day to a maximum recommended daily dose of 3000 mg. The effectiveness of doses lower than 3000 mg/day has not been adequately studied.

**Pediatric Patients Ages 6 To < 16 Years**  
 Treatment should be initiated with a daily dose of 20 mg/kg in 2 divided doses (10 mg/kg twice daily). The daily dose should be increased every 2 weeks by increments of 20 mg/kg to the recommended daily dose of 60 mg/kg (30 mg/kg twice daily). Psychiatric symptoms, behavioral abnormalities, suicidal thoughts/behavior, and/or unusual changes in mood or behavior (5.2).  
 Patients with body weight < 20 kg should be dosed on oral solution. Patients with body weight above 20 kg can be dosed with either tablets or oral solution (see Dosage and Administration (2.1)). Only white tablets above 20 kg are available.

**2.5 Adult Patients With Impaired Renal Function**  
 Levetiracetam dosing must be individualized according to the patient's renal function status. Recommended doses and adjustments are shown in Table 1. In order to calculate the dose, recommended oral patients with renal impairment, creatinine clearance adjusted for body surface area must be calculated. To this an estimate of the patient's creatinine clearance (CL<sub>CR</sub>) in mL/min must be factored using the following formula:

$$CL_{CR} = \frac{[140 - \text{years (males)}] \times \text{weight (kg)}}{72 \times \text{serum creatinine (mg/dL)}} \times 0.85 \text{ for female patients}$$

Then CL<sub>CR</sub> is adjusted for body surface area (BSA) as follows:

$$CL_{CR} (\text{mL/min/1.73m}^2) = \frac{CL_{CR} (\text{mL/min})}{BSA \text{ subject (m}^2)} \times 1.73$$

Table 1. Dosing Adjustment Regimen For Adult Patients With Impaired Renal Function

Group	Creatinine Clearance (mL/min/1.73m <sup>2</sup> )	Dosage (mg)	Frequency
Normal	> 80	500 to 1,500	Every 12 hours
Mild	50 - 80	500 to 1,000	Every 12 hours
Moderate	30 - 50	250 to 750	Every 12 hours
Severe	< 30	250 to 500	Every 12 hours
ESRD patients using dialysis	----	500 to 1,000 <sup>a</sup>	Every 24 hours <sup>b</sup>

<sup>a</sup> For pediatric patients: fatigue, aggression, nasal congestion, decreased appetite, and irritability (6.1)  
<sup>b</sup> To report SUSPECTED ADVERSE REACTIONS, contact Boca Pharmacal LLC, at 1-800-354-8468 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

----- **USE IN SPECIFIC POPULATIONS** -----  
 • Pregnancy: Plasma levels of levetiracetam may be decreased and therefore need to be monitored closely during pregnancy. Based on animal data, may cause fetal harm (5.9, 8.1)

**See 17 for Patient Counseling Information and Medication Guide.**

**5.3 REMOVAL OF ADVERSE REACTIONS**  
 Most common adverse reactions (incidence in levetiracetam-treated patients is ≥ 5% more than in placebo-treated patients) include:  
 • Adult patients: somnolence, asthenia, infection and dizziness (6.1)  
 • Pediatric patients: fatigue, aggression, nasal congestion, decreased appetite, and irritability (6.1)

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**5.5 Coordination Difficulties**

Coordination difficulties were only observed in the adult partial onset seizure studies. A total of 3.4% of adult levetiracetam-treated patients experienced coordination difficulties, (reported as either ataxia, abnormal gait, or incoordination) compared to 1.6% of placebo patients. A total of 0.4% of patients in controlled trials discontinued levetiracetam treatment due to ataxia, compared to 0.2% of placebo patients. In 0.7% of treated patients and in 0.2% of placebo patients the dose was reduced due to coordination difficulties, while one of the treated patients was hospitalized due to worsening of pre-existing ataxia. These events occurred most frequently within the first 4 weeks of treatment.

Patients should be monitored for these signs and symptoms and advised not to drive or operate machinery until they have gained sufficient experience on levetiracetam to gauge whether it could adversely affect their ability to drive or operate machinery.

**5.6 Withdrawal Seizures**  
 Antiepileptic drugs, including levetiracetam, should be withdrawn gradually to minimize the potential of increased seizure activity. Monitor patients for psychiatric signs and symptoms (5.1).

**5.7 Hematologic Abnormalities**  
**Partial Onset Seizures**  
 Adult patients statistically significant, decreased compared to placebo in total mean RBC count (0.03 × 10<sup>12</sup>/mL, mean hemoglobin (0.09 g/dL), and mean hematocrit (0.38%), were seen in levetiracetam-treated patients in controlled trials.

A total of 3.2% of treated and 1.8% of placebo patients had at least one possibly significant (<2.8 × 10<sup>12</sup>/L) decreased WBC, and 2.4% of treated and 1.4% of placebo patients had at least one possibly significant (<1.0 × 10<sup>9</sup>/L) decreased neutrophil count. Of the treated patients with a low neutrophil count, all but one case towards or to baseline with continued treatment. No patient was discontinued secondary to low neutrophil counts.

**Pediatric Patients 4 Years To < 16 Years**  
 Statistically significant decreases in WBC and neutrophil counts were seen in levetiracetam-treated patients as compared to placebo. The mean decreases from baseline in the levetiracetam-treated group were -0.4 × 10<sup>12</sup>/L and -0.3 × 10<sup>9</sup>/L, respectively, whereas there were small increases in the placebo group. Mean relative myelocyte counts increased by 1.7% in levetiracetam-treated patients, compared to a decrease of 4% in placebo patients (statistically significant).

In the controlled trial, more levetiracetam-treated patients had a possibly clinically significant abnormally low WBC value (3.0% levetiracetam-treated versus 0% placebo), however, there was no apparent difference between treatment groups with respect to neutrophil counts (0% levetiracetam-treated versus 4% placebo). No patient was discontinued secondary to low WBC or neutrophil counts.

In the controlled cognitive and neuropsychological safety study, two subjects (6.1%) in the placebo group and 5 subjects (8.6%) in the levetiracetam-treated group had high eosinophil count values that were possibly clinically significant (>10% or >0.7X10<sup>9</sup>/L).

**Juvenile Myoclonic Epilepsy**  
 Although there was no obvious hematologic abnormalities observed in patients with JME, the limited number of patients makes any conclusion tentative. The data from the partial seizure patients should be considered to be relevant for JME patients.

**5.8 Blood Pressure Increases**  
 In a randomized, placebo-controlled study in patients age 1 month to 4 years of age, a significantly higher risk of at least one measured increase in diastolic blood pressure was observed in the levetiracetam-treated patients (1.7%) compared to the placebo-treated patients (2%). There was no overall difference in mean diastolic blood pressure between the treatment groups. This disparity between the levetiracetam and placebo treatment groups was not observed in the study of older children.

**5.9 Seizure Control During Pregnancy**  
 Physiological changes may gradually decrease plasma levels of levetiracetam throughout pregnancy. This decrease was more pronounced during the third trimester. It is recommended that patients be monitored carefully during pregnancy. Close monitoring should continue through the postpartum period especially if the dose was changed during pregnancy.

**6.2 Description of Adverse Reactions**  
 The following adverse reactions are discussed in more details in other sections of labeling:

- Psychiatric Symptoms (see Warnings and Precautions (5.1))
- Suicidal Behavior and Ideation (see Warnings and Precautions (5.2))
- Somnolence and Fatigue (see Warnings and Precautions (5.3))
- Serious Dermatological Reactions (see Warnings and Precautions (5.4))
- Coordination Difficulties (see Warnings and Precautions (5.5))
- Withdrawal Seizures (see Warnings and Precautions (5.6))
- Hematologic Abnormalities (see Warnings and Precautions (5.7))
- Blood Pressure Increases (see Warnings and Precautions (5.8))
- Seizure Control During Pregnancy (see Warnings and Precautions (5.9))

**6.1 Clinical Trials Experience**  
 Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The prescriber should be aware that the adverse reaction incidence figures in the following tables, obtained when levetiracetam was added to concurrent AED therapy, cannot be used to predict the frequency of adverse reactions in the course of usual medical practice where patient characteristics and other factors may differ from those prevailing during clinical trials. Similarly, the cited frequencies cannot be directly compared with figures obtained from other controlled clinical trials involving different treatments, uses, or investigators. An inspection of these frequencies, however, does provide the prescriber with one basis to estimate the relative contribution of drug and non-drug factors to adverse reaction incidences in the population studied.

**Partial Onset Seizures**  
**Adults**  
 In controlled clinical studies in adults with partial onset seizures, the most frequently reported adverse reactions in patients receiving levetiracetam in combination with other AEDs were somnolence, asthenia, infection and dizziness. Of the most frequently reported adverse reactions in adults experiencing partial onset seizures, asthenia, somnolence and infection were reported with different frequencies. An inspection of these frequencies, however, does provide the prescriber with one basis to estimate the relative contribution of drug and non-drug factors to adverse reaction incidences in the population studied.

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The most common side effects seen in people who take levetiracetam tablets, USP include:

- sleepiness
- weakness
- infection
- dizziness

The most common side effects seen in children who take levetiracetam tablets, USP include, in addition to those listed above:

- tiredness
- acting aggressive
- nasal congestion
- decreased appetite
- irritability

These side effects can happen at any time but happen more often within the first 4 weeks of treatment except for infection.

Tell your healthcare provider if you have any side effect that bothers you or that does not go away.

These are not all the possible side effects of levetiracetam tablets, USP. For more information, ask your healthcare provider or pharmacist.

**Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.**

**How should I store levetiracetam tablets, USP?**

- Store levetiracetam tablets, USP at room temperature, 59°F to 86°F (15°C to 30°C) away from heat and light.
- **Keep levetiracetam tablets, USP and all medicines out of the reach of children.**

**General information about levetiracetam tablets, USP.**

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use levetiracetam tablets, USP for a condition for which it was not prescribed. Do not give levetiracetam tablets, USP to other people, even if they have the same symptoms that you have. It may harm them.

This Medication Guide summarizes the most important information about levetiracetam tablets, USP. If you would like more information, talk with your healthcare provider. You can ask your pharmacist or healthcare provider for information about levetiracetam tablets, USP that is written for health professionals.

**What are the ingredients of levetiracetam tablets, USP?**

**Levetiracetam tablets, USP** active ingredient: levetiracetam  
Inactive ingredients: colloidal silicon dioxide, corn starch, croscarmellose sodium, hypromellose, magnesium stearate, povidone, titanium dioxide, and additional agents listed below:

250 mg tablets: FD&C Blue #2 aluminum lake  
500 mg tablets: polyethylene glycol and yellow iron oxide  
750 mg tablets: FD&C Blue #2 aluminum lake, FD&C yellow #6 aluminum lake, polyethylene glycol and red iron oxide

Levetiracetam tablets, USP do not contain lactose or gluten.

*Information on the use of levetiracetam in pediatric patients less than 4 years of age as adjunctive therapy in the treatment of partial onset seizures is approved for UCB, Inc.'s levetiracetam tablets and oral solution. However, due to UCB, Inc.'s marketing exclusivity rights, this drug product is not labeled with that pediatric information.*

**Rx Only**

This Medication Guide has been approved by the US Food and Drug Administration.

Manufactured by: Cipla Ltd. Verna Goa, India

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Levetiracetam also displayed inhibitory properties in the kindling model in rats, another model of human complex partial seizures, both during kindling development and in the fully kindled state. The predictive value of these animal models for specific types of human epilepsy is uncertain.

*In vitro* and *in vivo* recordings of epileptiform activity from the hippocampus have shown that levetiracetam inhibits burst firing without affecting normal neuronal excitability, suggesting that levetiracetam may selectively prevent hypersynchronization of burst firing and propagation of seizure activity.

Levetiracetam at concentrations of up to 10 µM did not demonstrate binding affinity for a variety of known receptors, such as those associated with benzodiazepines, GABA (gamma-aminobutyric acid), glycine, NMDA (N-methyl-D-aspartate),  $\alpha$ -1-adrenergic, and second messenger systems. Furthermore, *in vitro* studies have failed to find an effect of levetiracetam on neuronal voltage-gated sodium or L-type calcium currents and levetiracetam does not appear to directly facilitate GABAergic neurotransmission. However, *in vitro* studies have demonstrated that levetiracetam opposes the activity of negative modulators of GABA- and glycine-gated currents and partially inhibits N-type calcium currents in neuronal cells.

A saturable and stereoselective neuronal binding site in rat brain tissue has been described for levetiracetam. Experimental data indicate that this binding site is the synaptic vesicle protein SV2A, thought to be involved in the regulation of vesicle exocytosis. Although the significance of levetiracetam binding to SV2A is not understood, levetiracetam and related analogs showed a rank order of affinity for SV2A which correlated with the potency of their antiseizure activity in autologous seizure-prone mice. These findings suggest that the interaction of levetiracetam with the SV2A protein may contribute to the antiepileptic mechanism of action of the drug.

**12.2 Pharmacokinetics**

**Effects on QTc Interval**

The effect of levetiracetam on QTc prolongation was evaluated in a randomized, double-blind, positive-controlled (moxifloxacin 400 mg) and placebo-controlled crossover study of levetiracetam (1000 mg or 5000 mg) in 52 healthy subjects. The upper bound of the 90% confidence interval for the largest placebo-adjusted, baseline-corrected QTc was below 10 milliseconds. Therefore, there was no evidence of significant QTc prolongation in this study.

**12.3 Pharmacokinetics**

**Absorption and Distribution**

Absorption of levetiracetam is rapid, with peak plasma concentrations occurring in about an hour following oral administration in fasted subjects. The oral bioavailability of levetiracetam tablets, USP is 100% and the tablets are bioequivalent in rate and extent of absorption to the oral solution. The elimination half-life of levetiracetam is 6.5 hours and decreases  $C_{max}$  by 20% and delays  $T_{max}$  by 1.5 hours. The pharmacokinetics of levetiracetam are linear over the dose range of 500-5000 mg. Steady state is achieved after 2 days of multiple twice-daily dosing. Levetiracetam and its major metabolite are less than 10% bound to plasma proteins; clinically significant interactions with other drugs through competition for protein binding sites are therefore unlikely.

**Metabolism**

Levetiracetam is not extensively metabolized in humans. The major metabolic pathway is the enzymatic hydrolysis of the acetamide group, which produces the carboxylic acid metabolite, ucb L057 (24% of dose) and is not dependent on any liver cytochrome P450 isoenzymes. The major metabolite is inactive in animal seizure models. Two minor metabolites were identified as the 2-oxo-pyrrolidine ring in position 5 (1% of dose). There is no enantiomeric interconversion of levetiracetam or its major metabolite.

**Elimination**

Levetiracetam plasma half-life in adults is 7 ± 1 hour and is unaffected by either dose or repeated administration. Levetiracetam is eliminated from the systemic circulation by renal excretion as unchanged drug which represents 66% of administered dose. The total body clearance is 0.96 mL/min/kg and the renal clearance is 0.6 mL/min/kg. The mechanism of excretion is glomerular filtration with subsequent partial tubular reabsorption. The metabolite ucb L057 is excreted by glomerular filtration and active tubular secretion with a renal clearance of 4 mL/min/kg. Levetiracetam elimination is correlated to creatinine clearance. Levetiracetam clearance is reduced in patients with impaired renal function (see *Use in Specific Populations (8.6) and Dosage and Administration (2.5)*).

**Specific Populations**

**Elderly**

Pharmacokinetics of levetiracetam were evaluated in 16 elderly subjects (age 61-88 years) with creatinine clearance ranging from 30 to 74 mL/min. Following oral administration of twice-daily dosing for 10 days, total body clearance decreased by 38% and the half-life was 2.5 hours longer in the elderly compared to healthy adults. This is most likely due to the decrease in renal function in these subjects.

**Pediatric Patients**

Pharmacokinetics of levetiracetam were evaluated in 24 pediatric patients (age 6-12 years) after single dose (20 mg/kg). The body weight adjusted apparent clearance of levetiracetam was approximately 40% higher than in adults.

A repeat dose pharmacokinetic study was conducted in pediatric patients (age 4-12 years) at doses of 20 mg/kg/day, 40 mg/kg/day, and 60 mg/kg/day. The evaluation of the pharmacokinetic profile of levetiracetam and its metabolite (ucb L057) in 14 pediatric patients demonstrated rapid absorption of levetiracetam at all doses with a  $T_{max}$  of about 1 hour and a  $t_{1/2}$  of 5 hours across the three dosing levels. The pharmacokinetics of levetiracetam in children was linear between 20 to 60 mg/kg/day. The potential interaction of levetiracetam with other AEDs was also evaluated in these patients. Levetiracetam had no significant effect on the plasma concentrations of carbamazepine, valproic acid, topiramate or lamotrigine. However, there was about a 22% increase of apparent clearance of levetiracetam when it was co-administered with an enzyme-inducing AED (e.g. carbamazepine).

**Pharmacokinetic information in pediatric patients less than 4 years of age is approved for UCB, Inc.'s levetiracetam tablets and oral solution. However, due to UCB, Inc.'s marketing exclusivity rights, this drug product is not labeled with that pediatric information.**

Population pharmacokinetic analysis showed that body weight was significantly correlated to the clearance of levetiracetam in pediatric patients; clearance increased with an increase in body weight.

**Pregnancy**

Levetiracetam levels may decrease during pregnancy.

**Gender**

Levetiracetam  $C_{max}$  and AUC were 20% higher in women (N=11) compared to men (N=12). However, clearances adjusted for body weight were comparable.

**Race**

Formal pharmacokinetic studies of the effects of race have not been conducted. Cross study comparisons involving Caucasians (N=12) and Asians (N=12), however, show that pharmacokinetics of levetiracetam were comparable between the two races. Because levetiracetam is primarily renally excreted and there are no important racial differences in creatinine clearance, pharmacokinetic differences due to race are not expected.

**Renal Impairment**

The disposition of levetiracetam was studied in adult subjects with varying degrees of renal function. Total body clearance of levetiracetam is reduced in patients with impaired renal function by 40% in the mild group (CL<sub>cr</sub> = 50-80 mL/min), 50% in the moderate group (CL<sub>cr</sub> = 30-50 mL/min) and 60% in the severe renal impairment group (CL<sub>cr</sub> <30 mL/min). Clearance of levetiracetam is correlated with creatinine clearance.

In anuric (end stage renal disease) patients, the total body clearance decreased 70% compared to normal subjects (CL<sub>cr</sub> >80 mL/min). Approximately 50% of the pool of levetiracetam in the body is removed during a standard 4-hour hemodialysis procedure.

Dosage should be reduced in patients with impaired renal function receiving levetiracetam, and supplemental doses should be given to patients after dialysis (see *Dosage and Administration (2.5)*).

**Hepatic Impairment**

In subjects with mild (Child-Pugh A) to moderate (Child-Pugh B) hepatic impairment, the pharmacokinetics of levetiracetam were unchanged. In patients with severe hepatic impairment (Child-Pugh C), total body clearance was 59% that of normal subjects, but decreased renal clearance accounted for most of the decrease. No dose adjustment is needed for patients with hepatic impairment.

**Drug Interactions**

*In vitro* data on metabolic interactions indicate that levetiracetam is unlikely to produce, or be subject to, pharmacokinetic interactions. Levetiracetam and its major metabolite, at concentrations well above  $C_{max}$  levels achieved within the therapeutic dose range, are neither inhibitors of, nor high affinity substrates for, human liver cytochrome P450 isoenzymes or UDP-glucuronidation enzymes. In addition, levetiracetam does not affect the *in vitro* glucuronidation of valproic acid.

Potential pharmacokinetic interactions of or with levetiracetam were assessed in clinical pharmacokinetic studies (phenytoin, valproate, warfarin, digoxin, oral contraceptive, probenecid) and through pharmacokinetic screening in the placebo-controlled clinical studies in epilepsy patients.

**Phenytoin**

Levetiracetam (3000 mg daily) had no effect on the pharmacokinetic disposition of phenytoin in patients with refractory epilepsy. Pharmacokinetics of levetiracetam were also not affected by phenytoin.

**Valproate**

Levetiracetam (1500 mg twice daily) did not alter the pharmacokinetics of valproate in healthy volunteers. Valproate 500 mg twice daily did not modify the rate or extent of levetiracetam absorption or its plasma clearance or urinary excretion. There also was no effect on exposure to and the excretion of the primary metabolite, ucb L057.

**Other Antiepileptic Drugs**

Potential drug interactions between levetiracetam and other AEDs (carbamazepine, gabapentin, lamotrigine, phenobarbital, phenytoin, valproate, warfarin, digoxin, oral contraceptive, probenecid) and through pharmacokinetic screening of levetiracetam and these AEDs during placebo-controlled clinical studies. These data indicate that levetiracetam does not influence the plasma concentration of other AEDs and that these AEDs do not influence the pharmacokinetics of levetiracetam.

**Effect of AEDs in Pediatric Patients**

There was about a 22% increase of apparent total body clearance of levetiracetam when it was co-administered with enzyme-inducing AEDs. Dose adjustment is not recommended. Levetiracetam had no effect on plasma concentrations of carbamazepine, valproate, topiramate, or lamotrigine.

**Oral Contraceptives**

Levetiracetam (500 mg twice daily) did not influence the pharmacokinetics of an oral contraceptive containing 0.03 mg ethinyl estradiol and 0.15 mg norgestrel, or of the estradiol hormone and progesterone levels, indicating that impairment of contraceptive efficacy is unlikely. Coadministration of this oral contraceptive did not influence the pharmacokinetics of levetiracetam.

**Digoxin**

Levetiracetam (1000 mg twice daily) did not influence the pharmacokinetics and pharmacodynamics (ECG) of digoxin given as a 0.25 mg dose every day. Coadministration of digoxin did not influence the pharmacokinetics of levetiracetam.

**Warfarin**

Levetiracetam (1000 mg twice daily) did not influence the pharmacokinetics of R and S warfarin. Prothrombin time was not affected by levetiracetam. Coadministration of warfarin did not affect the pharmacokinetics of levetiracetam.

**Probenecid**

Probenecid, a renal tubular secretion blocking agent, administered at a dose of 500 mg four times a day, did not change the pharmacokinetics of levetiracetam 1000 mg twice daily.  $C_{max}$  of the metabolite, ucb L057, was approximately doubled in the presence of probenecid while the fraction of drug excreted unchanged in the urine remained the same. Renal clearance of ucb L057 in the presence of probenecid decreased 60%, probably related to competitive inhibition of tubular secretion of ucb L057. The effect of levetiracetam on probenecid was not studied.

### 13 NONCLINICAL TOXICOLOGY

#### 13.1 Carcinogenesis, Mutagenesis, Impairment Of Fertility

**Carcinogenesis**

Rats were dosed with levetiracetam in the diet for 104 weeks at doses of 50, 300 and 1800 mg/kg/day. The highest dose is 6 times the maximum recommended daily human dose (MRHD) of 3000 mg on a mg/m<sup>2</sup> basis and also provided systemic exposure (AUC) approximately 6 times that achieved in humans receiving the MRHD. There was no evidence of

carcinogenicity. In mice, oral administration of levetiracetam for 80 weeks (doses up to 960 mg/kg/day) or 2 years (doses up to 4000 mg/kg/day, lowered to 3000 mg/kg/day after 45 weeks due to intertolerability) was not associated with an increase in tumors. The highest dose tested in mice for 2 years (3000 mg/kg/day) is approximately 5 times the MRHD on a mg/m<sup>2</sup> basis.

#### Mutagenesis

Levetiracetam was not mutagenic in the Ames test or in mammalian cells *in vitro* in the Chinese hamster ovary/HGPRT focus assay. It was not clastogenic in an *in vitro* analysis of metaphase chromosomes obtained from Chinese hamster ovary cells or in an *in vivo* mouse micronucleus assay. The hydrolysis product and major human metabolite of levetiracetam (ucb L057) was not mutagenic in the Ames test or the *in vitro* mouse lymphoma assay.

#### Impairment Of Fertility

No adverse effects on male or female fertility or reproductive performance were observed in rats at oral doses up to 1800 mg/kg/day (6 times the maximum recommended human dose on a mg/m<sup>2</sup> or systemic exposure (AUC) basis).

### 14 CLINICAL STUDIES

In the following studies, statistical significance versus placebo indicates a p value <0.05.

#### 14.1 Partial Onset Seizures

**Effectiveness in Partial Onset Seizures in Adults With Epilepsy**

The effectiveness of levetiracetam as adjunctive therapy (added to other antiepileptic drugs) in adults was established in three multicenter, randomized, double-blind, placebo-controlled clinical studies in patients who had refractory partial onset seizures with or without secondary generalization. The tablet formulation was used in all three studies. In these studies, 904 patients were randomized to placebo, 1000 mg, 2000 mg, or 3000 mg/day. Patients enrolled in Study 1 or Study 2 had refractory partial onset seizures for at least two years and had taken two or more classical AEDs. Patients enrolled in Study 3 had refractory partial onset seizures for at least 1 year and had taken one classical AED. At the time of the study, patients were taking a stable dose regimen of at least one and could take a maximum of two AEDs. During the baseline period, patients had to have experienced at least two partial onset seizures during each 4-week period.

**Study 1**

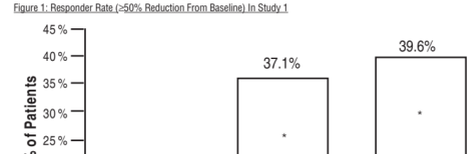
Study 1 was a double-blind, placebo-controlled, parallel-group study conducted at 41 sites in the United States comparing levetiracetam 1000 mg/day (N=97), levetiracetam 3000 mg/day (N=101), and placebo (N=95) given in equally divided doses twice daily. After a prospective baseline period of 12 weeks, patients were randomized to one of the three treatment groups described above. The 18-week treatment period consisted of a 6-week titration period, followed by a 12-week fixed dose evaluation period, during which concomitant AED regimens were held constant. The primary measure of effectiveness was a between group comparison of the percent reduction in weekly partial seizure frequency relative to placebo over the entire randomized treatment period (titration + evaluation period). Secondary outcome variables included the responder rate (incidence of patients with >50% reduction from baseline in partial onset seizure frequency). The results of the analysis of Study 1 are displayed in Table 10.

**Table 10. Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures in Study 1**

	Placebo (N=95)	Levetiracetam 1000 mg/day (N=97)	Levetiracetam 3000 mg/day (N=101)
Percent reduction in partial seizure frequency over placebo	-	26.1%*	30.1%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the three treatment groups (x-axis) is presented in Figure 1.



**Study 2**

Study 2 was a double-blind, placebo-controlled, crossover study conducted at 62 centers in Europe comparing levetiracetam 1000 mg/day (N=106), levetiracetam 2000 mg/day (N=105), and placebo (N=111) given in equally divided doses twice daily.

The first period of the study (Period A) was designed to be analyzed as a parallel-group study. After a prospective baseline period of up to 12 weeks, patients were randomized to one of the three treatment groups described above. The 16-week treatment period consisted of the 4-week titration period followed by a 12-week fixed dose evaluation period, during which concomitant AED regimens were held constant. The primary measure of effectiveness was a between group comparison of the percent reduction in weekly partial seizure frequency relative to placebo over the entire randomized treatment period (titration + evaluation period).

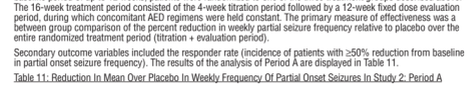
Secondary outcome variables included the responder rate (incidence of patients with >50% reduction from baseline in partial onset seizure frequency). The results of the analysis of Period A are displayed in Table 11.

**Table 11. Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures in Study 2: Period A**

	Placebo (N=111)	Levetiracetam 1000 mg/day (N=106)	Levetiracetam 2000 mg/day (N=101)
Percent reduction in partial seizure frequency over placebo	-	17.1%*	21.4%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the three treatment groups (x-axis) is presented in Figure 2.



\*statistically significant versus placebo

The comparison of levetiracetam 2000 mg/day to levetiracetam 1000 mg/day for responder rate was statistically significant (P=0.02). Analysis of the trial as a cross-over yielded similar results.

**Study 3**

Study 3 was a double-blind, placebo-controlled, parallel-group study conducted at 47 centers in Europe comparing levetiracetam 3000 mg/day (N=180) and placebo (N=104) in patients with refractory partial onset seizures, with or without secondary generalization, receiving only one concomitant AED. Study drug was given in two divided doses. After a 4-week baseline period of 12 weeks, patients were randomized to one of two treatment groups described above. The 16-week treatment period consisted of a 4-week titration period, followed by a 12-week fixed dose evaluation period, during which concomitant AED doses were held constant. The primary measure of effectiveness was a between group comparison of the percent reduction in weekly seizure frequency relative to placebo over the entire randomized treatment period (titration + evaluation period). Secondary outcome variables included the responder rate (incidence of patients with >50% reduction from baseline in partial onset seizure frequency). Table 12 displays the results of the analysis of Study 3.

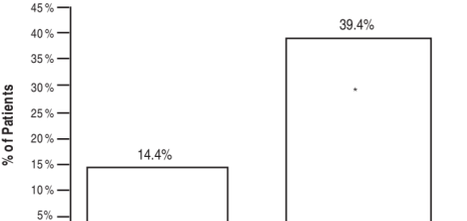
Table 12. Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures In Study 3

	Placebo (N=104)	Levetiracetam 3000 mg/day (N=180)
Percent reduction in partial seizure frequency over placebo	-	23.0%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) is presented in Figure 3.

**Figure 3. Responder Rate (>50% Reduction From Baseline) In Study 3**



\*statistically significant versus placebo

**Effectiveness in Partial Onset Seizures in Pediatric Patients 4 Years To 16 Years With Epilepsy**

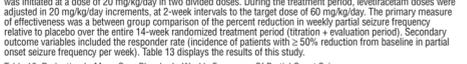
The effectiveness of levetiracetam as adjunctive therapy (added to other antiepileptic drugs) in pediatric patients was established in one multicenter, randomized double-blind, placebo-controlled study, conducted at 60 sites in North America, in children 4 to 16 years of age with partial seizures uncontrolled by standard antiepileptic drugs (AEDs). Eligible patients on a stable dose of 1-2 AEDs, who still experienced at least 4 partial onset seizures during the 4 weeks prior to screening, as well as at least 4 partial onset seizures in each of the two 4-week baseline periods, were randomized to receive either levetiracetam or placebo. The enrolled population included 198 patients (levetiracetam N=101, placebo N=97) with refractory partial onset seizures, whether or not secondarily generalized. The study consisted of an 8-week baseline period and a 2-week titration period followed by a 10-week evaluation period. Dosing was initiated at a dose of 20 mg/kg/day in two divided doses. During the treatment period, levetiracetam doses were adjusted in 20 mg/kg/day increments, at 2-week intervals to the target dose of 60 mg/kg/day. The primary measure of effectiveness was a between group comparison of the percent reduction in weekly partial seizure frequency relative to placebo over the entire 14-week randomized treatment period (titration + evaluation period). Secondary outcome variables included the responder rate (incidence of patients with >50% reduction from baseline in partial onset seizure frequency per week). Table 13 displays the results of this study.

**Table 13. Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures**

	Placebo (N=97)	Levetiracetam (N=101)
Percent reduction in partial seizure frequency over placebo	-	26.8%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) is presented in Figure 4.



**Table 13. Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures**

	Placebo (N=97)	Levetiracetam (N=101)
Percent reduction in partial seizure frequency over placebo	-	26.8%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) is presented in Figure 4.



\*statistically significant versus placebo

**Clinical trial information in pediatric patients less than 4 years of age as adjunctive therapy in the treatment of partial onset seizures is approved for UCB, Inc.'s levetiracetam tablets and oral solution. However, due to UCB, Inc.'s marketing exclusivity rights, this drug product is not labeled with that pediatric information.**

#### 14.2 Myoclonic Seizures in Patients With Juvenile Myoclonic Epilepsy

**Effectiveness Of Myoclonic Seizures in Patients >12 Years Of Age With Juvenile Myoclonic Epilepsy (JME)**

The effectiveness of levetiracetam as adjunctive therapy (added to other antiepileptic drugs) in patients 12 years of age and older with juvenile myoclonic epilepsy (JME) experiencing myoclonic seizures was established in one multicenter, randomized, double-blind, placebo-controlled study, conducted at 37 sites in 14 countries. Of the 120 patients enrolled, 113 had a diagnosis of confirmed or suspected JME. Eligible patients on a stable dose of 1 antiepileptic drug (AED) experiencing one or more myoclonic seizures per day for at least 8 days during the prospective 8-week baseline period were randomized to either levetiracetam or placebo (levetiracetam N=60, placebo N=60). Patients were treated over 4 weeks to a target dose of 3000 mg/day and treated at a stable dose of 3000 mg/day over 12 weeks (evaluation period). Study drug was given in 2 divided doses.

The primary measure of effectiveness was the proportion of patients with at least 50% reduction in the number of days per week with one or more myoclonic seizures during the treatment period (titration + evaluation periods) as compared to baseline. Table 14 displays the results for the 113 patients with JME in this study.

**Table 14. Responder Rate (>50% Reduction From Baseline) In Myoclonic Seizure Days Per Week For Patients With JME**

	Placebo (N=59)	Levetiracetam (N=54)
Percent of responders	23.7%	60.4%*

\*statistically significant versus placebo

**14.3 Primary Generalized Tonic-Clonic Seizures**

**Effectiveness in Primary Generalized Tonic-Clonic Seizures in Patients >6 Years Of Age**

The effectiveness of levetiracetam as adjunctive therapy (added to other antiepileptic drugs) in patients 6 years of age and older with idiopathic generalized epilepsy experiencing primary generalized tonic-clonic (PGTC) seizures was established in one multicenter, randomized, double-blind, placebo-controlled study, conducted at 50 sites in 8 countries. Eligible patients on a stable dose of 1 or 2 antiepileptic drugs (AEDs) experiencing at least 3 PGTC seizures during the 4-week combined baseline period and at least one PGTC seizure during the 4 weeks prior to the prospective baseline period and at least one PGTC seizure during the 4-week prospective baseline period) were randomized to either levetiracetam or placebo. The 8-week combined baseline period is referred to as "baseline" in the remainder of this section. The population included 164 patients (levetiracetam N=80, placebo N=84) with idiopathic generalized epilepsy (predominantly juvenile myoclonic epilepsy, juvenile absence epilepsy, childhood absence epilepsy, or epilepsy with Grand Mal seizures on awakening) experiencing primary generalized tonic-clonic seizures. Each of these syndromes of idiopathic generalized epilepsy was well represented in this patient population. Patients were treated over 4 weeks to a target dose of 3000 mg/day for adults or a pediatric target dose of 60 mg/kg/day and treated at a stable dose of 3000 mg/day (or 60 mg/kg/day for children) over 20 weeks (evaluation period). Study drug was given in 2 equally divided doses per day.

The primary measure of effectiveness was the percent reduction from baseline in weekly PGTC seizure frequency for levetiracetam and placebo treatment groups over the treatment period (titration + evaluation periods). There was a statistically significant decrease from baseline in PGTC seizure frequency in the levetiracetam-treated patients compared to the placebo-treated patients.

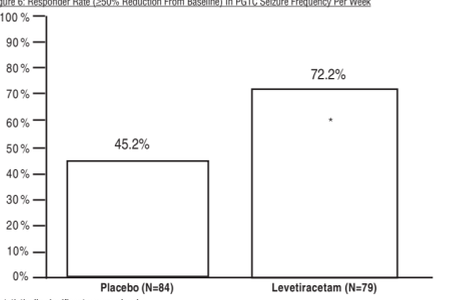
**Table 15. Median Percent Reduction From Baseline In PGTC Seizure Frequency Per Week**

	Placebo (N=84)	Levetiracetam (N=78)
Percent reduction in PGTC seizure frequency	44.8%	77.8%*

\*statistically significant versus placebo

The percentage of patients (y-axis) who achieved >50% reduction in weekly seizure rates from baseline in PGTC seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) is presented in Figure 6.

Figure 6. Responder Rate (>50% Reduction From Baseline) In PGTC Seizure Frequency Per Week



\*statistically significant versus placebo

**16 HOW SUPPLIED/STORAGE AND HANDLING**

**16.1 How Supplied**

Levetiracetam tablets, USP 250 mg are available as light blue, capsule-shaped, biconvex, film-coated tablets, debossed "750" and a score on one side and "0138" on the reverse side of the tablets, packaged in bottles of 60, 90, 120, and 500 tablets.

Levetiracetam tablets, USP 250 mg are available as follows:

- Bottles of 60 (NDC 64376-136-61)
- Bottles of 90 (NDC 64376-137-61)
- Bottles of 120 (NDC 64376-137-12)
- Bottles of 500 (NDC 64376-136-05)

Levetiracetam tablets,