HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use KEPPRA XR® safely and effectively. See full prescribing information for KEPPRA XR.

KEPPRA XR (levetiracetam) extended-release tablets, for oral use
Initial U.S. Approval: 1999

RECENT MAJOR CHANGES
Indications and Usage (1) [xx/2014]
Warnings and Precautions (5.1, 5.3, 5.7) [xx/2014]

INDICATIONS AND USAGE
KEPPRA XR is an antiepileptic drug indicated for adjunctive therapy in the treatment of partial onset seizures in patients ≥12 years of age with epilepsy (1)

DOSAGE AND ADMINISTRATION
KEPPRA XR is administered once daily; treatment should be initiated with a dose of 1000 mg once daily; the dosage may be adjusted in increments of 1000 mg every 2 weeks, to a maximum recommended dose of 3000 mg once daily (2.1)

KEPPRA XR dosing must be individualized according to renal function status (2.2)

DOSAGE FORMS AND STRENGTHS
• 500 mg white, film-coated extended-release tablet (3)
• 750 mg white, film-coated extended-release tablet (3)

CONTRAINDICATIONS
None (4)

WARNINGS AND PRECAUTIONS
Psychiatric Reactions: Monitor patients for behavioral abnormalities including psychotic symptoms, suicidal ideation, irritability, and aggressive behavior (5.1)
Suicidal Behavior and Ideation: Monitor patients for new or worsening depression, suicidal thoughts/behavior, and/or unusual changes in mood or behavior (5.2)
Somnolence and Fatigue: Monitor patients for these symptoms and advise patients not to drive or operate machinery until they have gained sufficient experience on KEPPRA XR (5.3)
Withdrawal Seizures: KEPPRA XR must be withdrawn gradually (5.6)

ADVERSE REACTIONS
Most common adverse reactions (incidence in KEPPRA XR–treated patients is ≥5% more than in placebo-treated patients) include somnolence and irritability (6.1)

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 (5.8, 8.1)

To report SUSPECTED ADVERSE REACTIONS, contact UCB, Inc. at 866-822-0068 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 (5.8, 8.1)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide

Revised: [08/2014]

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1 INDICATIONS AND USAGE

KEPPRA XR® is indicated as adjunctive therapy in the treatment of partial onset seizures in patients ≥12 years of age with epilepsy.

2 DOSAGE AND ADMINISTRATION

2.1 General Dosing Recommendations

KEPPRA XR is administered once daily. Treatment should be initiated with a dose of 1000 mg once daily. The once daily dosage may be adjusted in increments of 1000 mg every 2 weeks to a maximum recommended once daily dose of 3000 mg.

2.2 Adult Patients with Impaired Renal Function

KEPPRA XR dosing must be individualized according to the patient's renal function status. Recommended doses and adjustment for dose for adults are shown in Table 1. In order to calculate the dose recommended for patients with renal impairment, creatinine clearance adjusted for body surface area must be calculated. To do this, an estimate of the patient's creatinine clearance (CLcr) in mL/min must first be calculated using the following formula:

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

Then CLcr is adjusted for body surface area (BSA) as follows:

\[
CLcr (\text{mL/min}) = \frac{CLcr (\text{mL/min/1.73m}^2)}{\text{BSA subject (m}^2)} \times 1.73
\]

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

<table>
<thead>
<tr>
<th>Group</th>
<th>Creatinine Clearance (mL/min/1.73m²)</th>
<th>Dosage (mg)</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>&gt; 80</td>
<td>1000 to 3000</td>
<td>Every 24 hours</td>
</tr>
<tr>
<td>Mild</td>
<td>50 – 80</td>
<td>1000 to 2000</td>
<td>Every 24 hours</td>
</tr>
<tr>
<td>Moderate</td>
<td>30 – 50</td>
<td>500 to 1500</td>
<td>Every 24 hours</td>
</tr>
<tr>
<td>Severe</td>
<td>&lt; 30</td>
<td>500 to 1000</td>
<td>Every 24 hours</td>
</tr>
</tbody>
</table>

3 DOSAGE FORMS AND STRENGTHS

KEPPRA XR tablets are white, oblong-shaped, film-coated extended-release tablets imprinted in red with “UCB 500XR” on one side and contain 500 mg levetiracetam.

KEPPRA XR tablets are white, oblong-shaped, film-coated extended-release tablets imprinted in red with “UCB 750XR” on one side and contain 750 mg levetiracetam.

4 CONTRAINDICATIONS

None
5 WARNINGS AND PRECAUTIONS

5.1 Psychiatric Reactions

Patients on KEPPRA XR should be monitored for behavioral abnormalities.

KEPPRA XR Tablets

A total of 6.5% of patients treated with KEPPRA XR experienced non-psychotic behavioral disorders (reported as irritability and aggression), compared to no patients on placebo. Irritability was reported in 6.5% of patients treated with KEPPRA XR. Aggression was reported in 1.3% of patients treated with KEPPRA XR.

No patient discontinued treatment or had a dose reduction as a result of these adverse reactions.

There is considerably less controlled clinical trial experience with KEPPRA XR than with immediate-release KEPPRA tablets, and some adverse reactions observed with immediate-release KEPPRA may not have been detected in KEPPRA XR clinical trials because of limited number of patients. These adverse reactions may however occur in patients receiving KEPPRA XR.

Immediate-Release KEPPRA Tablets

A total of 13.3% of adult patients and 37.6% of pediatric patients (4 to 16 years of age) treated with immediate-release KEPPRA experienced non-psychotic behavioral symptoms (reported as aggression, agitation, anger, anxiety, apathy, depersonalization, depression, emotional lability, hostility, hyperkinesias, irritability, nervousness, neurosis, and personality disorder), compared to 6.2% and 18.6% of adult and pediatric patients on placebo. A randomized, double-blind, placebo-controlled study was performed to assess the neurocognitive and behavioral effects of immediate-release KEPPRA tablets as adjunctive therapy in pediatric patients (4 to 16 years of age). An exploratory analysis suggested a worsening in aggressive behavior in patients treated with immediate-release KEPPRA tablets in that study [see Use in Specific Populations (8.4)].

A total of 1.7% of adult patients treated with immediate-release KEPPRA discontinued treatment due to behavioral adverse events, compared to 0.2% of patients on placebo. The treatment dose was reduced in 0.8% of adult patients treated with immediate-release KEPPRA, compared to 0.5% of patients on placebo. Overall, 10.9% of pediatric patients treated with immediate-release KEPPRA experienced behavioral symptoms associated with discontinuation or dose reduction, compared to 6.2% of pediatric patients on placebo.

One percent of adult patients and 2% of pediatric patients (4 to 16 years of age) treated with immediate-release KEPPRA experienced psychotic symptoms, compared to 0.2% and 2%, respectively, in adult and pediatric patients on placebo. In the controlled study that assessed the neurocognitive and behavioral effects of immediate-release KEPPRA in pediatric patients 4 to 16 years of age, 1 (1.6%) patient treated with KEPPRA experienced paranoia, compared to no patients on placebo. There were 2 (3.1%) patients treated with immediate-release KEPPRA who experienced confusional state, compared to no patients on placebo [see Use in Specific Populations (8.4)].

Two (0.3%) adult patients treated with immediate-release KEPPRA were hospitalized, and their treatment was discontinued due to psychosis. In both patients, the psychosis event developed within the first week of treatment, and resolved within 1 to 2 weeks following treatment discontinuation. There was no difference between drug and placebo-treated patients in the incidence of pediatric patients who discontinued treatment due to psychotic and non-psychotic adverse reactions.

5.2 Suicidal Behavior and Ideation

Antiepileptic drugs (AEDs), including KEPPRA XR, increase the risk of suicidal thoughts or behavior in patients taking these drugs for any indication. Patients treated with any AED for any indication should be monitored for the emergence or worsening of depression, suicidal thoughts or behavior, and/or any unusual changes in mood or behavior.

Pooled analyses of 199 placebo-controlled clinical trials (mono- and adjunctive therapy) of 11 different AEDs showed that patients randomized to one of the AEDs had approximately twice the risk (adjusted Relative Risk 1.8, 95% CI:1.2, 2.7) of suicidal thinking or behavior compared to patients randomized to placebo. In these trials, which had a median treatment duration of 12 weeks, the estimated incidence rate of suicidal behavior or ideation among 27,863 AED-treated patients was 0.43%, compared to 0.24% among 16,029 placebo-treated patients, representing an increase of approximately one case of suicidal thinking or behavior for every 530 patients treated. There were four suicides in drug-treated patients in the trials and none in placebo-treated patients, but the number is too small to allow any conclusion about drug effect on suicide.
The increased risk of suicidal thoughts or behavior with AEDs was observed as early as one week after starting drug treatment with AEDs and persisted for the duration of treatment assessed. Because most trials included in the analysis did not extend beyond 24 weeks, the risk of suicidal thoughts or behavior beyond 24 weeks could not be assessed.

The risk of suicidal thoughts or behavior was generally consistent among drugs in the data analyzed. The finding of increased risk with AEDs of varying mechanisms of action and across a range of indications suggests that the risk applies to all AEDs used for any indication. The risk did not vary substantially by age (5-100 years) in the clinical trials analyzed. Table 2 shows absolute and relative risk by indication for all evaluated AEDs.

### Table 2: Risk By Indication For Antiepileptic Drugs In The Pooled Analysis

<table>
<thead>
<tr>
<th>Indication</th>
<th>Placebo Patients with Events Per 1000 Patients</th>
<th>Drug Patients with Events Per 1000 Patients</th>
<th>Relative Risk: Incidence of Events in Drug Patients/Incidence in Placebo Patients</th>
<th>Risk Difference: Additional Drug Patients with Events Per 1000 Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epilepsy</td>
<td>1.0</td>
<td>3.4</td>
<td>3.5</td>
<td>2.4</td>
</tr>
<tr>
<td>Psychiatric</td>
<td>5.7</td>
<td>8.5</td>
<td>1.5</td>
<td>2.9</td>
</tr>
<tr>
<td>Other</td>
<td>1.0</td>
<td>1.8</td>
<td>1.9</td>
<td>0.9</td>
</tr>
<tr>
<td>Total</td>
<td>2.4</td>
<td>4.3</td>
<td>1.8</td>
<td>1.9</td>
</tr>
</tbody>
</table>

The relative risk for suicidal thoughts or behavior was higher in clinical trials for epilepsy than in clinical trials for psychiatric or other conditions, but the absolute risk differences were similar for the epilepsy and psychiatric indications.

Anyone considering prescribing KEPPRA XR or any other AED must balance the risk of suicidal thoughts or behavior with the risk of untreated illness. Epilepsy and many other illnesses for which AEDs are prescribed are themselves associated with morbidity and mortality and an increased risk of suicidal thoughts and behavior. Should suicidal thoughts and behavior emerge during treatment, the prescriber needs to consider whether the emergence of these symptoms in any given patient may be related to the illness being treated.

Patients, their caregivers, and families should be informed that AEDs increase the risk of suicidal thoughts and behavior and should be advised of the need to be alert for the emergence or worsening of the signs and symptoms of depression, any unusual changes in mood or behavior, or the emergence of suicidal thoughts, behavior, or thoughts about self-harm. Behaviors of concern should be reported immediately to healthcare providers.

### 5.3 Somnolence and Fatigue

Patients should be monitored for somnolence and fatigue, and be advised not to drive or operate machinery until they have gained sufficient experience on KEPPRA XR to gauge whether it adversely affects their ability to drive or operate machinery. In clinical trials of immediate-release KEPPRA, somnolence and asthenia occurred most frequently within the first 4 weeks of treatment.

#### KEPPRA XR Tablets

In the KEPPRA XR double-blind, controlled trial in patients experiencing partial onset seizures, 7.8% of patients treated with KEPPRA XR experienced somnolence, compared to 2.5% of patients on placebo.

No patient discontinued treatment or had a dose reduction as a result of these adverse reactions.

There is considerably less controlled clinical trial experience with KEPPRA XR than with immediate-release KEPPRA tablets, and some adverse reactions observed with immediate-release KEPPRA may not have been detected in KEPPRA XR clinical trials because of limited number of patients. These adverse reactions may however occur in patients receiving KEPPRA XR.

#### Immediate-Release KEPPRA Tablets

In general, the incidences of somnolence and fatigue in the pediatric partial onset seizure studies were comparable to those of the adult partial onset seizure studies.
In controlled trials of adult patients with epilepsy experiencing partial onset seizures, 14.8% of patients treated with immediate-release KEPPRA reported somnolence, compared to 8.4% of patients on placebo. There was no clear dose response up to 3000 mg/day. In a study in which there was no titration, about 45% of patients receiving 4000 mg/day reported somnolence. The somnolence was considered serious in 0.3% of patients treated with immediate-release KEPPRA, compared to no patients on placebo. About 3% of patients treated with immediate-release KEPPRA discontinued treatment due to somnolence, compared to 0.7% of patients on placebo. The dose was reduced due to somnolence in 1.4% of patients treated with immediate-release KEPPRA and in 0.9% of patients on placebo, while 0.3% of the patients treated with immediate-release KEPPRA were hospitalized due to somnolence.

In controlled trials of adult patients with epilepsy experiencing partial onset seizures, 14.7% of patients treated with immediate-release KEPPRA reported asthenia, compared to 9.1% of patients on placebo. Treatment was discontinued due to asthenia in 0.8% of patients treated with immediate-release KEPPRA, compared to 0.5% of patients on placebo. The dose was reduced due to asthenia in 0.5% of patients treated with immediate-release KEPPRA and in 0.2% of patients on placebo.

5.4 Serious Dermatological Reactions
Serious dermatological reactions, including Stevens-Johnson syndrome (SJS) and toxic epidermal necrolysis (TEN), have been reported in both children and adults treated with KEPPRA. The median time of onset is reported to be 14 to 17 days, but cases have been reported at least four months after initiation of treatment. Recurrence of the serious skin reactions following rechallenge with KEPPRA has also been reported. KEPPRA XR should be discontinued at the first sign of a rash, unless the rash is clearly not drug-related. If signs or symptoms suggest SJS/TEN, use of this drug should not be resumed and alternative therapy should be considered.

5.5 Coordination Difficulties
Patients should be monitored for coordination difficulties and advised not to drive or operate machinery until they have gained sufficient experience on KEPPRA XR to gauge whether it could adversely affect their ability to drive or operate machinery.

A total of 3.4% of adult patients treated with immediate-release KEPPRA experienced coordination difficulties, (reported as either ataxia, abnormal gait, or incoordination), compared to 1.6% of patients on placebo. A total of 0.4% of patients in controlled trials discontinued immediate-release KEPPRA treatment due to ataxia, compared to no patients on placebo. In 0.7% of patients treated with immediate-release KEPPRA and in 0.2% of patients on placebo, 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.

Coordination difficulties were not observed in the KEPPRA XR controlled trial. There is considerably less controlled clinical trial experience with KEPPRA XR than with immediate-release KEPPRA tablets, and some adverse reactions observed with immediate-release KEPPRA may not have been detected in KEPPRA XR clinical trials because of limited number of patients. These adverse reactions may however occur in patients receiving KEPPRA XR.

5.6 Withdrawal Seizures
Antiepileptic drugs, including KEPPRA XR, should be withdrawn gradually to minimize the potential of increased seizure frequency.

5.7 Hematologic Abnormalities
Although there were no obvious hematologic abnormalities observed in the KEPPRA XR controlled study, there is considerably less controlled clinical trial experience with KEPPRA XR than with immediate-release KEPPRA tablets, and some adverse reactions observed with immediate-release KEPPRA may not have been detected in KEPPRA XR clinical trials because of limited number of patients. These adverse reactions may however occur in patients receiving KEPPRA XR.

In controlled trials, a minor but statistically significant decrease (compared to placebo) in total mean RBC count (0.03 x 10^6/mm^3), mean hemoglobin (0.09 g/dL), and mean hematocrit (0.38%), was seen in adult patients treated with immediate-release KEPPRA. A total of 3.2% of adult patients treated with immediate-release KEPPRA, and 1.8% of patients on placebo had at least one possibly significant (≤2.8 x 10^9/L) decreased WBC, and 2.4% of patients treated with immediate-release KEPPRA vs. 1.4% of patients on placebo had at least one possibly significant (≤1.0 x 10^9/L) decreased neutrophil count. Of the patients treated with immediate-release KEPPRA with a low neutrophil count, all but one rose towards or to baseline with continued treatment. No patient was discontinued secondary to low neutrophil counts.
In pediatric patients (4 to <16 years of age), statistically significant decreases in WBC and neutrophil counts were seen in patients treated with immediate-release KEPPRA, as compared to placebo. The mean decreases from baseline in the immediate-release KEPPRA group were \(-0.4 \times 10^9/L\) and \(-0.3 \times 10^9/L\), respectively, whereas there were small increases in the placebo group. A significant increase in mean relative lymphocyte counts was observed in 1.7% of patients treated with immediate-release KEPPRA compared to a decrease of 4% in patients on placebo.

In the controlled pediatric trial, a possibly clinically significant abnormal low WBC value was observed in 3% of patients treated with immediate-release KEPPRA, compared to no patients on placebo. However, there was no apparent difference between treatment groups with respect to neutrophil count. No patient was discontinued secondary to low WBC or neutrophil counts.

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

5.8 Seizure Control During Pregnancy
Physiological changes may gradually decrease plasma levels of levetiracetam throughout pregnancy. This decrease is 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 ADVERSE REACTIONS
The following adverse reactions are discussed in more details in other sections of labeling:

- Psychiatric Reactions [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)]

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 table, obtained when KEPPRA XR 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 clinical investigations 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 the adverse reaction incidences in the population studied.

KEPPRA XR Tablets
In the controlled clinical study using KEPPRA XR in patients with partial onset seizures (Study 1), the most frequently reported adverse reactions in patients receiving KEPPRA XR in combination with other AEDs, for events with rates greater than placebo, were irritability and somnolence.

Table 3 lists adverse reactions that occurred in at least 5% of epilepsy patients treated with KEPPRA XR participating in the placebo-controlled study (Study 1) and were numerically more common than in patients treated with placebo. In this study, either KEPPRA XR or placebo was added to concurrent AED therapy. Adverse reactions were usually mild to moderate in intensity.

<table>
<thead>
<tr>
<th>Body System/ Adverse Reaction</th>
<th>KEPPRA XR (N=77) %</th>
<th>Placebo (N=79) %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastrointestinal Disorders</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 4 lists the adverse reactions seen in the controlled studies of immediate-release KEPPRA tablets in adult patients experiencing partial onset seizures. Although the pattern of adverse reactions in the KEPPRA XR study seems somewhat different from that seen in partial onset seizure controlled studies for immediate-release KEPPRA tablets, this is possibly due to the much smaller number of patients in this study compared to the immediate-release tablet studies. The adverse reactions for KEPPRA XR are expected to be similar to those seen with immediate-release KEPPRA tablets.

**Immediate-Release KEPPRA Tablets**

**Adults**

In controlled clinical studies of immediate-release KEPPRA tablets as adjunctive therapy to other AEDs in adults with partial onset seizures, the most frequently reported adverse reactions, for events with rates greater than placebo, were somnolence, asthenia, infection and dizziness.

Table 4 lists adverse reactions that occurred in at least 1% of adult epilepsy patients treated with immediate-release KEPPRA tablets participating in placebo-controlled studies and were numerically more common than in patients treated with placebo. In these studies, either immediate-release KEPPRA tablets or placebo was added to concurrent AED therapy. Adverse reactions were usually mild to moderate in intensity.

**Table 4: Incidence (%) of Adverse Reactions in Placebo-Controlled, Add-On Studies in Adults Experiencing Partial Onset Seizures By Body System (Adverse Reactions Occurred in At Least 1% of Immediate-Release KEPPRA-Treated Patients and Occurred More Frequently Than Placebo-Treated Patients)**
**Pediatric Patients 4 Years to <16 Years**

In a pooled analysis of two controlled pediatric clinical studies in children 4 to 16 years of age with partial onset seizures, the adverse reactions most frequently reported with the use of immediate-release KEPPRA in combination with other AEDs, and with greater frequency than in patients on placebo, were fatigue, aggression, nasal congestion, decreased appetite, and irritability.

Table 5 lists adverse reactions that occurred in at least 2% of pediatric patients treated with immediate-release KEPPRA and were more common than in pediatric patients on placebo. In these studies, either immediate-release KEPPRA or placebo was added to concurrent AED therapy. Adverse reactions were usually mild to moderate in intensity.

Table 5: Incidence (%) Of Adverse Reactions In Pooled Placebo-Controlled, Add-On Studies In Pediatric Patients Ages 4 to 16 Years Experiencing Partial Onset Seizures By Body System (Adverse Reactions Occurred In At Least 2% Of Patients Treated With Immediate-Release KEPPRA And Occurred More Frequently Than Patients on Placebo)
<table>
<thead>
<tr>
<th>Body System/Adverse Reaction</th>
<th>Immediate-Release KEPPRA (N=165) %</th>
<th>Placebo (N=131) %</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ear and Labyrinth Disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ear Pain</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><strong>Eye Disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conjunctivitis</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><strong>Gastrointestinal Disorders</strong></td>
<td></td>
<td></td>
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<tr>
<td>Vomiting</td>
<td>15</td>
<td>12</td>
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<tr>
<td>Abdominal Pain Upper</td>
<td>9</td>
<td>8</td>
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<td>Diarrhea</td>
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<td>Constipation</td>
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<td><strong>General Disorders and Administration Site Conditions</strong></td>
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<td>Fatigue</td>
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<td><strong>Infections and Infestations</strong></td>
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<td>Nasopharyngitis</td>
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<td>Influenza</td>
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<td>1</td>
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<tr>
<td>Gastroenteritis</td>
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<td>0</td>
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<td>Rhinitis</td>
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<td>0</td>
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<tr>
<td><strong>Injury, Poisoning and Procedural Complications</strong></td>
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<td>Head Injury</td>
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<td>Contusion</td>
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<td>Fall</td>
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<td>Joint Sprain</td>
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<td><strong>Metabolism and Nutrition Disorders</strong></td>
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<tr>
<td>Decreased Appetite</td>
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<td>Anorexia</td>
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<td><strong>Musculoskeletal and Connective Tissue Disorders</strong></td>
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<td>Arthralgia</td>
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<td><strong>Nervous System</strong></td>
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<tr>
<td>Headache</td>
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<tr>
<td>Somnolence</td>
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<td>9</td>
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<tr>
<td>Dizziness</td>
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Reference ID: 3603477
Cough | 9 | 5
Nasal Congestion | 9 | 2
Pharyngolaryngeal Pain | 7 | 4

In controlled pediatric clinical studies in patients 4-16 years of age, 7% of patients treated with immediate-release KEPPRA tablets and 9% of patients on placebo discontinued as a result of an adverse event.

In addition, the following adverse reactions were seen in other well-controlled studies of immediate-release KEPPRA tablets: balance disorder, disturbance in attention, eczema, hyperkinesia, memory impairment, myalgia, personality disorders, pruritus, and vision blurred.

Comparison of Gender, Age and Race
There are insufficient data for KEPPRA XR to support a statement regarding the distribution of adverse experience reports by gender, age and race.

6.2 Postmarketing Experience
In addition to the adverse reactions listed above for immediate-release KEPPRA tablets [see Adverse Reactions (6.1)], the following adverse reactions have been identified during post-approval use of immediate-release KEPPRA tablets. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure. The listing is alphabetized: abnormal liver function test, choreoathetosis, dyskinesia, erythema multiforme, hepatic failure, hepatitis, leukopenia, muscular weakness, neutropenia, pancreatitis, pancytopenia (with bone marrow suppression identified in some of these cases), panic attack, thrombocytopenia, and weight loss. Alopecia has been reported with immediate-release KEPPRA use; recovery was observed in majority of cases where immediate-release KEPPRA was discontinued.

7 DRUG INTERACTIONS

No significant pharmacokinetic interactions were observed between levetiracetam or its major metabolite and concomitant medications via human liver cytochrome P450 isofoms, epoxide hydrolase, UDP-glucuronidation enzymes, P-glycoprotein, or renal tubular secretion [see Clinical Pharmacology (12.3)].

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy
KEPPRA XR levels may decrease during pregnancy [see Warnings and Precautions (5.8)].

Pregnancy Category C
There are no adequate and well-controlled studies in pregnant women. In animal studies, levetiracetam produced evidence of developmental toxicity, including teratogenic effects, at doses similar to or greater than human therapeutic doses. KEPPRA XR should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Oral administration of levetiracetam to female rats throughout pregnancy and lactation led to increased incidences of minor fetal skeletal abnormalities and retarded offspring growth pre- and/or postnatally at doses ≥350 mg/kg/day (equivalent to the maximum recommended human dose of 3000 mg [MRHD] on a mg/m² basis) and with increased pup mortality and offspring behavioral alterations at a dose of 1800 mg/kg/day (6 times the MRHD on a mg/m² basis). The developmental no effect dose was 70 mg/kg/day (0.2 times the MRHD on a mg/m² basis). There was no overt maternal toxicity at the doses used in this study.

Oral administration of levetiracetam to female rats throughout pregnancy and lactation led to increased incidences of minor fetal skeletal abnormalities and retarded offspring growth pre- and/or postnatally at doses ≥350 mg/kg/day (equivalent to the maximum recommended human dose of 3000 mg [MRHD] on a mg/m² basis) and with increased pup mortality and offspring behavioral alterations at a dose of 1800 mg/kg/day (6 times the MRHD on a mg/m² basis). The developmental no effect dose was 70 mg/kg/day (0.2 times the MRHD on a mg/m² basis). There was no overt maternal toxicity at the doses used in this study.

Oral administration of levetiracetam to pregnant rabbits during the period of organogenesis resulted in increased embryofetal mortality and increased incidences of minor fetal skeletal abnormalities at doses ≥600 mg/kg/day (4 times MRHD on a mg/m² basis) and in decreased fetal weights and increased incidences of fetal malformations at a dose of 1800 mg/kg/day (12 times the MRHD on a mg/m² basis). The developmental no effect dose was 200 mg/kg/day (equivalent to the MRHD on a mg/m² basis). Maternal toxicity was also observed at 1800 mg/kg/day.

When levetiracetam was administered orally to pregnant rats during the period of organogenesis, fetal weights were decreased and the incidence of fetal skeletal variations was increased at a dose of 3600 mg/kg/day (12 times the MRHD). 1200 mg/kg/day (4 times the MRHD) was a developmental no effect dose. There was no evidence of maternal toxicity in this study.
Treatment of rats with levetiracetam during the last third of gestation and throughout lactation produced no adverse developmental or maternal effects at oral doses of up to 1800 mg/kg/day (6 times the MRHD on a mg/m² basis).

Pregnancy Registries
To provide information regarding the effects of in utero exposure to KEPPRA XR, physicians are advised to recommend that pregnant patients taking KEPPRA XR enroll in the North American Antiepileptic Drug (NAAED) pregnancy registry. This can be done by calling the toll free number 1-888-233-2334, and must be done by the patients themselves. Information on the registry can also be found at the website http://www.aedpregnancyregistry.org/.

UCB, Inc. has established the UCB AED Pregnancy Registry to advance scientific knowledge about safety and outcomes in pregnant women being treated with all UCB antiepileptic drugs including KEPPRA XR. To ensure broad program access and reach, either a healthcare provider or the patient can initiate enrollment in the UCB AED Pregnancy Registry by calling (888) 537-7734 (toll free).

8.2 Labor and Delivery
The effect of KEPPRA XR on labor and delivery in humans is unknown.

8.3 Nursing Mothers
Levetiracetam is excreted in human milk. Because of the potential for serious adverse reactions in nursing infants from KEPPRA XR, a decision should be made whether to discontinue nursing or discontinue the drug, taking into account the importance of the drug to the mother.

8.4 Pediatric Use
Safety and effectiveness in pediatric patients 12 years of age and older has been established based on pharmacokinetic data in adults and adolescents using KEPPRA XR and efficacy and safety data in controlled pediatric studies using immediate-release KEPPRA [see Adverse Reactions (6.1), Clinical Pharmacology (12.3), and Clinical Studies (14.1)].

Safety and effectiveness of KEPPRA XR in patients below the age of 12 years have not been established.

A 3-month, randomized, double-blind, placebo-controlled study was performed to assess the neurocognitive and behavioral effects of immediate-release KEPPRA as adjunctive therapy in 98 pediatric patients with inadequately controlled partial seizures, ages 4 to 16 years (KEPPRA N=64; placebo N=34). The target dose of immediate-release KEPPRA was 60 mg/kg/day. Neurocognitive effects were measured by the Leiter-R Attention and Memory (AM) Battery, which assesses various aspects of a child's memory and attention. Although no substantive differences were observed between the placebo- and KEPPRA-treated groups in the median change from baseline in this battery, the study was not adequate to assess formal statistical non-inferiority between the drug and placebo. The Achenbach Child Behavior Checklist (CBCL/6-18), a standardized validated tool used to assess a child’s competencies and behavioral/emotional problems, was also assessed in this study. An analysis of the CBCL/6-18 indicated a worsening in aggressive behavior, one of the eight syndrome scores, in patients treated with KEPPRA [see Warnings and Precautions (5.1)].

Studies of levetiracetam in juvenile rats (dosing from day 4 through day 52 of age) and dogs (dosing from week 3 through week 7 of age) at doses of up to 1800 mg/kg/day (approximately 7 and 24 times, respectively, the maximum recommended pediatric dose of 60 mg/kg/day on a mg/m² basis) did not indicate a potential for age-specific toxicity.

8.5 Geriatric Use
There were insufficient numbers of elderly subjects in controlled trials of epilepsy to adequately assess the effectiveness of KEPPRA XR in these patients. It is expected that the safety of KEPPRA XR in elderly patients 65 and over would be comparable to the safety observed in clinical studies of immediate-release KEPPRA tablets.

There were 347 subjects in clinical studies of immediate-release levetiracetam that were 65 and over. No overall differences in safety were observed between these subjects and younger subjects. There were insufficient numbers of elderly subjects in controlled trials of epilepsy to adequately assess the effectiveness of immediate-release KEPPRA in these patients.

Levetiracetam is known to be substantially excreted by the kidney, and the risk of adverse reactions to this drug may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function [see Clinical Pharmacology (12.3)].
8.6 Use in Patients with Impaired Renal Function
The effect of KEPPRA XR on renally impaired patients was not assessed in the controlled study. However, it is expected that the effect on KEPPRA XR-treated patients would be similar to the effect seen in controlled studies of immediate-release KEPPRA tablets. Clearance of levetiracetam is decreased in patients with renal impairment and is correlated with creatinine clearance [see Clinical Pharmacology (12.3)]. Dose adjustment is recommended for patients with impaired renal function [see Dosage and Administration (2.1)].

10 OVERDOSAGE

10.1 Signs, Symptoms and Laboratory Findings of Acute Overdosage in Humans
The signs and symptoms for KEPPRA XR overdose are expected to be similar to those seen with immediate-release KEPPRA tablets.

The highest known dose of oral immediate-release KEPPRA received in the clinical development program was 6000 mg/day. Other than drowsiness, there were no adverse reactions in the few known cases of overdose in clinical trials. Cases of somnolence, agitation, aggression, depressed level of consciousness, respiratory depression and coma were observed with immediate-release KEPPRA overdoses in postmarketing use.

10.2 Management of Overdose
There is no specific antidote for overdose with KEPPRA XR. If indicated, elimination of unabsorbed drug should be attempted by emesis or gastric lavage; usual precautions should be observed to maintain airway. General supportive care of the patient is indicated including monitoring of vital signs and observation of the patient’s clinical status. A Certified Poison Control Center should be contacted for up to date information on the management of overdose with KEPPRA XR.

10.3 Hemodialysis
Standard hemodialysis procedures result in significant clearance of levetiracetam (approximately 50% in 4 hours) and should be considered in cases of overdose. Although hemodialysis has not been performed in the few known cases of overdose, it may be indicated by the patient's clinical state or in patients with significant renal impairment.

11 DESCRIPTION
KEPPRA XR is an antiepileptic drug available as 500 mg and 750 mg (white) extended-release tablets for oral administration.

The chemical name of levetiracetam, a single enantiomer, is (S)-α-ethyl-2-oxo-1-pyrrolidine acetamide, its molecular formula is C₈H₁₄N₂O₂ and its molecular weight is 170.21. Levetiracetam is chemically unrelated to existing antiepileptic drugs (AEDs). It has the following structural formula:

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CH₃CH₂C\(\text{CONH}_2\)
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Levetiracetam is a white to off-white crystalline powder with a faint odor and a bitter taste. It is very soluble in water (104.0 g/100 mL). It is freely soluble in chloroform (65.3 g/100 mL) and in methanol (53.6 g/100 mL), soluble in ethanol (16.5 g/100 mL), sparingly soluble in acetonitrile (5.7 g/100 mL) and practically insoluble in n-hexane. (Solubility limits are expressed as g/100 mL solvent.)

KEPPRA XR tablets contain the labeled amount of levetiracetam. Inactive ingredients: colloidal anhydrous silica, hypromellose, magnesium stearate, polyethylene glycol 6000, polyvinyl alcohol-partially hydrolyzed, titanium dioxide (E171), Macrogol/PEG3350, and talc. The imprinting ink contains shellac, FD&C Red #40, n-butyl alcohol, propylene glycol, titanium dioxide, ethanol, and methanol.

12 CLINICAL PHARMACOLOGY
12.1 Mechanism of Action

The precise mechanism(s) by which levetiracetam exerts its antiepileptic effect is unknown. The antiepileptic activity of levetiracetam was assessed in a number of animal models of epileptic seizures. Levetiracetam did not inhibit single seizures induced by maximal stimulation with electrical current or different chemoconvulsants and showed only minimal activity in submaximal stimulation and in threshold tests. Protection was observed, however, against secondarily generalized activity from focal seizures induced by pilocarpine and kainic acid, two chemoconvulsants that induce seizures that mimic some features of human complex partial seizures with secondary generalization. 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 epileptiform 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), re-uptake sites, and second messenger systems. Furthermore, in vitro studies have failed to find an effect of levetiracetam on neuronal voltage-gated sodium or T-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 molecular significance of levetiracetam binding to synaptic vesicle protein 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 audiogenic 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 Pharmacodynamics

Effects on QTc Interval

The effects of KEPPRA XR on QTc prolongation is expected to be the same as that of immediate-release KEPPRA. The effect of immediate-release KEPPRA on QTc prolongation was evaluated in a randomized, double-blind, positive-controlled (moxifloxacin 400 mg) and placebo-controlled crossover study of immediate-release KEPPRA (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

Overview

Bioavailability of KEPPRA XR tablets is similar to that of the immediate-release KEPPRA Tablets. The pharmacokinetics (AUC and C_{max}) were shown to be dose proportional after single dose administration of 1000 mg, 2000 mg, and 3000 mg extended-release levetiracetam. Plasma half-life of extended-release levetiracetam is approximately 7 hours.

Levetiracetam is almost completely absorbed after oral administration. The pharmacokinetics of levetiracetam are linear and time-invariant, with low intra- and inter-subject variability. Levetiracetam is not significantly protein-bound (<10% bound) and its volume of distribution is close to the volume of intracellular and extracellular water. Sixty-six percent (66%) of the dose is renally excreted unchanged. The major metabolic pathway of levetiracetam (24% of dose) is an enzymatic hydrolysis of the acetamide group. It is not liver cytochrome P450 dependent. The metabolites have no known pharmacological activity and are renally excreted. Plasma half-life of levetiracetam across studies is approximately 6-8 hours. The half-life is increased in the elderly (primarily due to impaired renal clearance) and in subjects with renal impairment.

Absorption and Distribution

Extended-release levetiracetam peak plasma concentrations occur in about 4 hours. The time to peak plasma concentrations is about 3 hours longer with extended-release levetiracetam than with immediate-release tablets.
Single administration of two 500 mg extended-release levetiracetam tablets once daily produced comparable maximal plasma concentrations and area under the plasma concentration versus time as did the administration of one 500 mg immediate-release tablet twice daily in fasting conditions. After multiple dose extended-release levetiracetam tablets intake, extent of exposure (AUC0-24) was similar to extent of exposure after multiple dose immediate-release tablets intake. Cmax and Cmin were lower by 17% and 26% after multiple dose extended-release levetiracetam tablets intake in comparison to multiple dose immediate-release tablets intake. Intake of a high fat, high calorie breakfast before the administration of extended-release levetiracetam tablets resulted in a higher peak concentration, and longer median time to peak. The median time to peak (Tmax) was 2 hours longer in the fed state.

Two 750 mg extended-release levetiracetam tablets were bioequivalent to a single administration of three 500 mg extended-release levetiracetam tablets.

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 product of hydroxylation of the 2-oxo-pyrrolidine ring (2% of dose) and opening of 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 (2.1)].

Special Populations
Elderly
There are insufficient pharmacokinetic data to specifically address the use of extended-release levetiracetam in the elderly population.

Pharmacokinetics of immediate-release 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
An open label, multicenter, parallel-group, two-arm study was conducted to evaluate the pharmacokinetics of KEPPRA XR in pediatric patients (13 to 16 years old) and in adults (18 to 55 years old) with epilepsy. KEPPRA XR oral tablets (1000 mg to 3000 mg) were administered once daily with a minimum of 4 days and a maximum of 7 days of treatment to 12 pediatric patients and 13 adults in the study. Dose-normalized steady-state exposure parameters, Cmax and AUC, were comparable between pediatric and adult patients.

Pregnancy
KEPPRA XR levels may decrease during pregnancy.

Gender
Extended-release levetiracetam Cmax was 21-30% higher and AUC was 8-18% higher in women (N=12) 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 with extended-release or immediate-release levetiracetam. Cross study comparisons involving Caucasians (N=12) and Asians (N=12), however, show that pharmacokinetics of immediate-release 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 effect of KEPPRA XR on renally impaired patients was not assessed in the controlled study. However, it is expected that the effect on KEPPRA XR-treated patients would be similar to that seen in controlled studies of immediate-release KEPPRA tablets. In patients with end stage renal disease on dialysis, it is recommended that immediate-release KEPPRA be used instead of KEPPRA XR.

The disposition of immediate-release 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 (CLcr = 50-80 mL/min), 50% in the moderate group (CLcr = 30-50 mL/min) and 60% in the severe renal impairment group (CLcr <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 (CLcr >80mL/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; immediate-release levetiracetam should be given to patients on dialysis [see Dosage and Administration (2.1)].

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 50% 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 isoforms, epoxide hydrolase 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 with immediate-release KEPPRA tablets in the placebo-controlled clinical studies in epilepsy patients. The potential for drug interactions for KEPPRA XR is expected to be essentially the same as that with immediate-release KEPPRA tablets.

Phenytoin
Immediate-release KEPPRA tablets (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
Immediate-release KEPPRA tablets (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 immediate-release KEPPRA tablets and other AEDs (carbamazepine, gabapentin, lamotrigine, phenobarbital, phenytoin, primidone and valproate) were also assessed by evaluating the serum concentrations 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.

Oral Contraceptives
Immediate-release KEPPRA tablets (500 mg twice daily) did not influence the pharmacokinetics of an oral contraceptive containing 0.03 mg ethinyl estradiol and 0.15 mg levonorgestrel, or of the luteinizing hormone and progesterone levels, indicating that impairment of contraceptive efficacies is unlikely. Coadministration of this oral contraceptive did not influence the pharmacokinetics of levetiracetam.
Digoxin
Immediate-release KEPPRA tablets (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
Immediate-release KEPPRA tablets (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\text{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 immediate-release KEPPRA tablets 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/m² basis and it 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 intolerability) 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/m² basis.

Mutagenesis
Levetiracetam was not mutagenic in the Ames test or in mammalian cells in vitro in the Chinese hamster ovary/HGPRT locus 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/m² or systemic exposure [AUC] basis).

14 CLINICAL STUDIES

The effectiveness of KEPPRA XR as adjunctive therapy in partial onset seizures in adults was established in one multicenter, randomized, double-blind, placebo-controlled clinical study in patients who had refractory partial onset seizures with or without secondary generalization. This was supported by the demonstration of efficacy of immediate-release KEPPRA tablets (see below) in partial seizures in three multicenter, randomized, double-blind, placebo-controlled clinical studies in adults, as well as a demonstration of comparable bioavailability between the XR and immediate-release formulations [see Clinical Pharmacology (12.3)] in adults. The effectiveness for KEPPRA XR as adjunctive therapy in partial onset seizures in pediatric patients, 12 years of age and older, was based upon a single pharmacokinetic study showing comparable pharmacokinetics of KEPPRA XR in adults and adolescents [see Clinical Pharmacology (12.3)]. All studies are described below.

14.1 KEPPRA XR in Adults
The effectiveness of KEPPRA XR as adjunctive therapy (added to other antiepileptic drugs) was established in one multicenter, randomized, double-blind, placebo-controlled clinical study across 7 countries in patients who had refractory partial onset seizures with or without secondary generalization (Study 1).

Study 1
Patients enrolled in Study 1 had at least eight partial seizures with or without secondary generalization during the 8-week baseline period and at least two partial seizures in each 4-week interval of the baseline period. Patients were taking a stable dose regimen of at least one AED, and could take a maximum of three AEDs. After a prospective baseline period of 8 weeks, 158 patients were randomized to placebo (N=79) or 1000 mg (two 500 mg tablets) of KEPPRA XR (N=79), given once daily over a 12-week treatment period.

The primary efficacy endpoint in Study 1 was the percent reduction over placebo in mean weekly frequency of partial onset seizures. The median percent reduction in weekly partial onset seizure frequency from baseline over the treatment period was 46.1% in the KEPPRA XR 1000 mg treatment group (N=74) and 33.4% in the placebo group (N=78). The estimated percent reduction over placebo in weekly partial onset seizure frequency over the treatment period was 14.4% (statistically significant).

The relationship between the effectiveness of the same daily dose of KEPPRA XR and immediate-release KEPPRA has not been studied and is unknown.

14.2 Immediate-release KEPPRA in Adults
The effectiveness of immediate-release KEPPRA 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 (Studies 2, 3, and 4). The tablet formulation was used in all three studies. In these studies, 904 patients were randomized to placebo, KEPPRA 1000 mg, KEPPRA 2000 mg, or KEPPRA 3000 mg/day. Patients enrolled in Study 2 or Study 3 had refractory partial onset seizures for at least two years, and had taken two or more AEDs. Patients enrolled in Study 4 had refractory partial onset seizures for at least 1 year and had taken one AED. At the time of the study, patients were taking a stable dose regimen of at least one AED, 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 2
Study 2 was a double-blind, placebo-controlled, parallel-group study conducted at 41 sites in the United States, comparing immediate-release KEPPRA 1000 mg/day (N=97), immediate-release KEPPRA 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 in Study 2 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 in Study 2 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 Study 2 are displayed in Table 6.

Table 6: Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures In Study 2

<table>
<thead>
<tr>
<th></th>
<th>Placebo (N=95)</th>
<th>Immediate-release KEPPRA 1000 mg/day (N=97)</th>
<th>Immediate-release KEPPRA 3000 mg/day (N=101)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent reduction in partial seizure frequency over placebo</td>
<td>–</td>
<td>26.1%*</td>
<td>30.1%*</td>
</tr>
</tbody>
</table>

* statistically significant versus placebo

The percentage of patients (y-axis) who achieved ≥50% reduction from baseline in weekly partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the three treatment groups (x-axis) in Study 2 is presented in Figure 1.
Study 3

Study 3 was a double-blind, placebo-controlled, crossover study conducted at 62 centers in Europe, comparing immediate-release KEPPRA 1000 mg/day (N=106), immediate-release KEPPRA 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 in Study 3 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 in Study 3 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 7.

### Table 7: Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures In Study 3: Period A

<table>
<thead>
<tr>
<th></th>
<th>Placebo (N=111)</th>
<th>Immediate-release KEPPRA 1000 mg/day (N=106)</th>
<th>Immediate-release KEPPRA 2000 mg/day (N=105)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent reduction in</td>
<td>–</td>
<td>17.1%*</td>
<td>21.4%*</td>
</tr>
<tr>
<td>partial seizure frequency over placebo</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* statistically significant versus placebo

The percentage of patients (y-axis) who achieved ≥50% reduction from baseline in weekly partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the three treatment groups (x-axis) in Study 3 is presented in Figure 2.
The comparison of immediate-release KEPPRA 2000 mg/day to immediate-release KEPPRA 1000 mg/day for responder rate in Study 3 was statistically significant ($P=0.02$). Analysis of the trial as a cross-over study yielded similar results.

**Study 4**

Study 4 was a double-blind, placebo-controlled, parallel-group study conducted at 47 centers in Europe comparing immediate-release KEPPRA 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 prospective baseline period of 12 weeks, patients in Study 4 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 in Study 4 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 $\geq 50\%$ reduction from baseline in partial onset seizure frequency). Table 8 displays the results of Study 4.

**Table 8: Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures In Study 4**

<table>
<thead>
<tr>
<th></th>
<th>Placebo ($N=104$)</th>
<th>Immediate-release KEPPRA 3000 mg/day ($N=180$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent reduction in partial seizure frequency over placebo</td>
<td>$-$</td>
<td>23.0%*</td>
</tr>
</tbody>
</table>

* statistically significant versus placebo

The percentage of patients (y-axis) who achieved $\geq 50\%$ reduction from baseline in weekly partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) in Study 4 is presented in Figure 3.
Figure 3: Responder Rate (≥50% Reduction From Baseline) In Study 4

![Graph showing responder rate comparison between Placebo (N=97) and Immediate-release KEPPRA 3000 mg/day (N=180).]

* statistically significant versus placebo

14.3 Immediate-release KEPPRA in Pediatric Patients 4 Years to 16 Years

The use of KEPPRA XR in pediatric patients 12 years of age and older is supported by Study 5, which was conducted using immediate-release KEPPRA. KEPPRA XR is not indicated in children below 12 years of age.

Study 5

The effectiveness of immediate-release KEPPRA as adjunctive therapy in pediatric patients was established in a 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 (Study 5). 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 immediate-release KEPPRA or placebo. The enrolled population included 198 patients (KEPPRA N=101; placebo N=97) with refractory partial onset seizures, with or without secondarily generalization. Study 5 consisted of an 8-week baseline period and 4-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, the immediate-release KEPPRA 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 in Study 5 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 9 displays the results of this study.

Table 9: Reduction In Mean Over Placebo In Weekly Frequency Of Partial Onset Seizures in Study 5

<table>
<thead>
<tr>
<th></th>
<th>Placebo (N=97)</th>
<th>Immediate-release KEPPRA (N=101)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percent reduction in partial seizure frequency over placebo</td>
<td>-</td>
<td>26.8%*</td>
</tr>
</tbody>
</table>

*statistically significant versus placebo

The percentage of patients (y-axis) who achieved ≥ 50% reduction in weekly partial onset seizure frequency over the entire randomized treatment period (titration + evaluation period) within the two treatment groups (x-axis) in Study 5 is presented in Figure 4.

Figure 4: Responder Rate (≥ 50% Reduction From Baseline) in Study 5
16 HOW SUPPLIED/STORAGE AND HANDLING

16.1 How Supplied

KEPPRA XR 500 mg tablets are white, oblong-shaped, film-coated tablets imprinted with “UCB 500XR” in red on one side. They are supplied in white HDPE bottles containing 60 tablets (NDC 50474-598-66).

KEPPRA XR 750 mg tablets are white, oblong-shaped, film-coated tablets imprinted with “UCB 750XR” in red on one side. They are supplied in white HDPE bottles containing 60 tablets (NDC 50474-599-66).

16.2 Storage

Store at 25°C (77°F); excursions permitted to 15-30°C (59-86°F) [see USP Controlled Room Temperature].

17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA-approved Patient Labeling (Medication Guide).

Counsel patients on the benefits and risks of receiving KEPPRA XR. Provide the Medication Guide to patients and/or caregivers, and instruct them to read the Medication Guide prior to taking KEPPRA XR. Instruct patients to take KEPPRA XR only as prescribed.

Suicidal Behavior and Ideation

Counsel patients, their caregivers, and/or families that antiepileptic drugs (AEDs), including KEPPRA XR, may increase the risk of suicidal thoughts and behavior and advise patients to be alert for the emergence or worsening of symptoms of depression; unusual changes in mood or behavior; or suicidal thoughts, behavior, or thoughts about self-harm. Advise patients, their caregivers, and/or families to immediately report behaviors of concern to a healthcare provider.

Psychiatric Reactions and Changes in Behavior

Advise patients that KEPPRA XR may cause changes in behavior (e.g. irritability and aggression). In addition, patients should be advised that they may experience changes in behavior that have been seen with other formulations of KEPPRA, which include agitation, anger, anxiety, apathy, depression, hostility, and, in rare cases, psychotic symptoms.

Effects on Driving or Operating Machinery

Inform patients that KEPPRA XR may cause dizziness and somnolence. Inform patients not to drive or operate machinery until they have gained sufficient experience on KEPPRA XR to gauge whether it adversely affects their ability to drive or operate machinery.

*statistically significant versus placebo
Dermatological Adverse Reactions
Advise patients that serious dermatological adverse reactions have occurred in patients treated with levetiracetam and instruct them to call their physician immediately if a rash develops.

Dosing and Administration
Patients should be instructed to only take KEPPRA XR once daily and to swallow the tablets whole. They should not be chewed, broken, or crushed.

Pregnancy
Advise patients to notify their healthcare provider if they become pregnant or intend to become pregnant during KEPPRA XR therapy. Encourage patients to enroll in the North American Antiepileptic Drug (NAAED) pregnancy registry if they become pregnant. This registry is collecting information about the safety of antiepileptic drugs during pregnancy. To enroll, patients can call the toll free number 1-888-233-2334. Additionally, inform patients they can enroll in the UCB AED Pregnancy Registry and they or their healthcare provider can call 1-888-537-7734 (toll free) [see Use In Specific Populations (8.1)].
MEDICATION GUIDE

KEPPRA XR® (KEPP-ruh XR) (levetiracetam)
extended-release tablets

Read this Medication Guide before you start taking KEPPRA XR and each time you get a refill. There may be new information. This information does not take the place of talking to your healthcare provider about your medical condition or treatment.

What is the most important information I should know about KEPPRA XR?

Like other antiepileptic drugs, KEPPRA XR may cause suicidal thoughts or actions in a very small number of people, about 1 in 500 people taking it.

Call a healthcare provider right away if you have any of these symptoms, especially if they are new, worse, or worry you:

- thoughts about suicide or dying
- attempts to commit suicide
- new or worse depression
- new or worse anxiety
- feeling agitated or restless
- panic attacks
- trouble sleeping (insomnia)
- new or worse irritability
- acting aggressive, being angry, or violent
- acting on dangerous impulses
- an extreme increase in activity and talking (mania)
- other unusual changes in behavior or mood

Do not stop KEPPRA XR without first talking to a healthcare provider.

- Stopping KEPPRA XR suddenly can cause serious problems. Stopping a seizure medicine suddenly can cause seizures that will not stop (status epilepticus).

- Suicidal thoughts or actions can be caused by things other than medicines. If you have suicidal thoughts or actions, your healthcare provider may check for other causes.

How can I watch for early symptoms of suicidal thoughts and actions?

- Pay attention to any changes, especially sudden changes, in mood, behaviors, thoughts, or feelings.

- Keep all follow-up visits with your healthcare provider as scheduled.
Call your healthcare provider between visits as needed, especially if you are worried about symptoms.

**What is KEPPRA XR?**

KEPPRA XR is a prescription medicine taken by mouth that is used with other medicines to treat partial onset seizures in people 12 years of age and older with epilepsy.

It is not known if KEPPRA XR is safe or effective in people under 12 years of age.

Before taking your medicine, make sure you have received the correct medicine. Compare the name above with the name on your bottle and the appearance of your medicine with the description of KEPPRA XR provided below. Tell your pharmacist immediately if you think you have been given the wrong medicine.

500 mg KEPPRA XR tablets are white, oblong-shaped, film-coated tablets marked with “UCB 500XR” in red on one side.

750 mg KEPPRA XR tablets are white, oblong-shaped, film-coated tablets marked with “UCB 750XR” in red on one side.

**What should I tell my healthcare provider before starting KEPPRA XR?**

Before taking KEPPRA XR, tell your healthcare provider about all of your medical conditions, including if you:

- have or have had depression, mood problems or suicidal thoughts or behavior
- have kidney problems
- are pregnant or planning to become pregnant. It is not known if KEPPRA XR will harm your unborn baby. You and your healthcare provider will have to decide if you should take KEPPRA XR while you are pregnant. If you become pregnant while taking KEPPRA XR, talk to your healthcare provider about registering with the North American Antiepileptic Drug Pregnancy Registry. You can enroll in this registry by calling 1-888-233-2334. You can also enroll in the UCB AED Pregnancy Registry by calling 1-888-537-7734. The purpose of these registries is to collect information about the safety of KEPPRA XR and other antiepileptic medicine during pregnancy.
- are breast feeding. KEPPRA XR can pass into your milk and may harm your baby. You and your healthcare provider should discuss whether you should take KEPPRA XR or breast-feed; you should not do both.

Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements. Do not start a new medicine without first talking with your healthcare provider.
Know the medicines you take. Keep a list of them to show your healthcare provider and pharmacist each time you get a new medicine.

**How should I take KEPPRA XR?**

Take KEPPRA XR exactly as prescribed.

- Your healthcare provider will tell you how much KEPPRA XR to take and when to take it. KEPPRA XR is usually taken once a day. Take KEPPRA XR at the same time each day.
- Your healthcare provider may change your dose. Do not change your dose without talking to your healthcare provider.
- Take KEPPRA XR with or without food.
- Swallow the tablets whole. Do not chew, break, or crush tablets.
- If you miss a dose of KEPPRA XR, take it as soon as you remember. If it is almost time for your next dose, just skip the missed dose. Take the next dose at your regular time. **Do not take two doses at the same time.**
- If you take too much KEPPRA XR, call your local Poison Control Center or go to the nearest emergency room right away.

**What should I avoid while taking KEPPRA XR?**

Do not drive, operate machinery or do other dangerous activities until you know how KEPPRA XR affects you. KEPPRA XR may make you dizzy or sleepy.

**What are the possible side effects of KEPPRA XR?**

- See “What is the most important information I should know about KEPPRA XR?”

KEPPRA XR can cause serious side effects.

Call your healthcare provider right away if you have any of these symptoms:

- mood and behavior changes such as aggression, agitation, anger, anxiety, apathy, mood swings, depression, hostility, and irritability. A few people may get psychotic symptoms such as hallucinations (seeing or hearing things that are really not there), delusions (false or strange thoughts or beliefs) and unusual behavior.
- extreme sleepiness, tiredness, and weakness
- problems with muscle coordination (problems walking and moving)
- a skin rash. Serious skin rashes can happen after you start taking KEPPRA XR. There is no way to tell if a mild rash will become a serious reaction.

Common side effects seen in people who take KEPPRA XR and other formulations of KEPPRA include:

- sleepiness

Reference ID: 3603477
• weakness
• infection
• dizziness

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 KEPPRA XR. For more information, ask your healthcare provider or pharmacist.

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

How should I store KEPPRA XR?

• Store KEPPRA XR at room temperature, 59°F to 86°F (15°C to 30°C) away from heat and light.
• Keep KEPPRA XR and all medicines out of the reach of children.

General information about the safe and effective use of KEPPRA XR.

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use KEPPRA XR for a condition for which it was not prescribed. Do not give KEPPRA XR 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 KEPPRA XR. If you would like more information, talk with your healthcare provider. You can ask your pharmacist or healthcare provider for information about KEPPRA XR that is written for health professionals. You can also get information about KEPPRA XR at www.keppraxr.com or call 1-866-822-0068.

What are the ingredients of KEPPRA XR?

**KEPPRA XR tablet** active ingredient: levetiracetam
Inactive ingredients: colloidal anhydrous silica, hypromellose, magnesium stearate, polyethylene glycol 6000, polyvinyl alcohol-partially hydrolyzed, titanium dioxide (E171), Macrogol/PEG3350, and talc. The imprinting ink contains shellac, FD&C Red #40, n-butyl alcohol, propylene glycol, titanium dioxide, ethanol, and methanol.

KEPPRA XR does not contain lactose or gluten.