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CNS SPECTRUMS THE INTERNATIONAL JOURNAL OF NEUROPSYCHIATRIC MEDICINE

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Because I don't want to lose my son to the voices again.

The voices in his head are back. I can't bear to see him like this.

He was doing so well on his own. This will ruin everything. It could send him back to the hospital.

We're fighting to get things back under control. But we need help now.



For resources to help you help your patients with schizophrenia, visit www.ToolsForTheFight.com

The labeling for ZYPREXA includes a boxed warning:

- Elderly patients with dementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk of death compared to placebo.
- ZYPREXA is not approved for the treatment of elderly patients with dementia-related psychosis.

ZYPREXA is approved for the treatment of schizophrenia, acute bipolar mania, and for maintenance treatment in bipolar disorder.

For Important Safety Information, including boxed warning, see adjacent page and accompanying Brief Summary of Prescribing Information.

Lilly

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Increased Mortality in Elderly Patients with Dementia-Related Psychosis

Elderly patients with dementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk of death compared to placebo. Analyses of seventeen placebo-controlled trials (modal duration of 10 weeks) in these patients revealed a risk of death in the drug-treated patients of between 1.6 to 1.7 times that seen in placebo-treated patients. Over the course of a typical 10-week controlled trial, the rate of death in drug-treated patients was about 4.5%, compared to a rate of about 2.6% in the placebo group. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular (eg, heart failure, sudden death) or infectious (eg, pneumonia) in nature. ZYPREXA is not approved for the treatment of elderly patients with dementia-related psychosis.

Cerebrovascular adverse events (CVAE), including stroke, in elderly patients with dementia—Cerebrovascular adverse events (eg, stroke, transient ischemic attack), including fatalities, were reported in patients in trials of ZYPREXA in elderly patients with dementia-related psychosis. In placebo-controlled trials, there was a significantly higher incidence of CVAE in patients treated with ZYPREXA compared to patients treated with placebo. ZYPREXA is not approved for the treatment of patients with dementia-related psychosis.

Hyperglycemia and diabetes mellitus—Hyperglycemia, in some cases associated with ketoacidosis, coma, or death, has been reported in patients treated with atypical antipsychotics including ZYPREXA. All patients taking atypicals should be monitored for symptoms of hyperglycemia. Persons with diabetes who are started on atypicals should be monitored regularly for worsening of glucose control; those with risk factors for diabetes should undergo baseline and periodic fasting blood glucose testing. Patients who develop symptoms of hyperglycemia during treatment should undergo fasting blood glucose testing.

Neuroleptic malignant syndrome (NMS)—As with all antipsychotic medications, a rare and potentially fatal condition known as NMS has been reported with olanzapine. If signs and symptoms appear, immediate discontinuation is recommended. Clinical manifestations of NMS are hyperpyrexia, muscle rigidity, altered mental status and evidence of autonomic instability (irregular pulse or blood pressure, tachycardia, diaphoresis and cardiac dysrhythmia). Additional signs may include elevated creatinine phosphokinase, myoglobinuria (rhabdomyolysis), and acute renal failure.

Tardive dyskinesia (TD)—As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of TD. The risk of developing TD and the likelihood that it will become irreversible are believed to increase as the duration of treatment and the total cumulative dose of antipsychotic increase. The syndrome may remit, partially or completely, if antipsychotic treatment is withdrawn.

Medication dispensing and prescribing errors have occurred between ZYPREXA® (olanzapine) and Zyrtec® (cetirizine HCl). These errors could result in unnecessary adverse events or potential relapse in patients suffering from schizophrenia or bipolar disorder. To reduce the potential for dispensing errors, please write ZYPREXA clearly.

The most common treatment-emergent adverse event associated with ZYPREXA (vs placebo) in 6-week acute-phase schizophrenia trials was somnolence (26% vs 15%). Other common events were dizziness (11% vs 4%), weight gain (6% vs 1%), personality disorder (COSTART term for nonaggressive objectionable behavior; 8% vs 4%), constipation (9% vs 3%), akathisia (5% vs 1%), and postural hypotension (5% vs 2%).

The most common treatment-emergent adverse event associated with ZYPREXA (vs placebo) in 3- and 4-week bipolar mania trials was somnolence (35% vs 13%). Other common events were dry mouth (22% vs 7%), dizziness (18% vs 6%), asthenia (15% vs 6%), constipation (11% vs 5%), dyspepsia (11% vs 5%), increased appetite (6% vs 3%), and tremor (6% vs 3%).

For complete safety profile, see the full Prescribing Information.

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ZYPREXA® (Olanzapine Tablets) ZYPREXA® ZYDIS® (Olanzapine Orally Disintegrating Tablets) ZYPREXA® IntraMuscular (Olanzapine for Injection) Brief Summary: Please consult package insert for complete prescribing information.

WARNING

WARNING increased Mortality in Elderly Patients with Dementia-Related Psychosis—Elderly patients with dementia-related psychosis treated with atypical antipsycholic drugs are at an increased risk of death compared to placebo. Analyses of seventeen placebo-controlled trials (modal duration of 10 weeks) in these patients revealed a risk of death in the drug-treated patients of between 1.6 to 1.7 times that seen in placebo-treated patients. Over the course of a typical 10-week controlled trial, the rate of death in drug-treated patients was about 4.5%, compared to a rate of about 2.6% in the placebo group. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular (eg, heart failure, sudden death) or infectious (eg, pneumonia) in nature. ZYPREXA is not approved for the treatment of patients with dementia-related psychosis.

INDICATIONS AND USAGE: ZYPREXA and ZYPREXA Zydis are indicated for short- and long-term treatment of schizophrenia, for acute manic and mixed episodes of bipolar I disorder, and for maintenance treatment in bipolar disorder. The use of ZYPREXA for extended periods should be periodically re-evaluated as to the long-term usefulness of the drug for the individual patient. ZYPREXA IntraMuscular is indicated for treatment of agitation associated with schizophrenia and bipolar I mania.

CONTRAINDICATIONS: Known hypersensitivity to olanzapine.

WARNINGS: <u>increased Mortality in Elderly Patients with Dementia-Related Psychosis</u>—Elderly patients with dementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk of death compared to placebo. ZYPREXA is not approved for the treatment of patients with dementia-related psychosis (see BOX WARNING).

compared to placebo. ZYPREXA is not approved for the treatment of patients with dementia-related psychosis (see BOX WARNING). In placebo-controlled clinical trials of elderly patients with dementia-related psychosis, the incidence of death in ofanzapine-treated patients (3.5%) was significantly greater than placebo-controlled trials. Ts%). <u>Carebrovascular Adverse Events. Including Stroke, in Elderly Patients with Dementia</u>—Cerebrovascular adverse events (eg. stroke, transient ischemic attack), including fatalities, were reported in patients in trials of olanzapine in elderly patients with dementia-related psychosis. In placebo-controlled trials, there was a significantly trighter incidence of cerebrovascular adverse events in patients treated with ofanzapine compared to patients treated with placebo. Olanzapine is not approved for the treatment of patients with dementia-related psychosis. <u>Hypercylowenia and Diabetes Mellitus</u>—Hypergyloemia, in some cases associated with ketoacidosis, coma, or death, has been reported in patients treated with atypical antipsychotics including olanzapine. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes in patients diagnosed with diabetes who are started on atypical antipsychotics should be monifored regularly for worsening of glucose control. Patients with risk factors for diabetes who are starting treatment with atypicals should have fasting blood glucose (FBG) testing at baseline and periodically during treatment. Any patient treated with atypical antipsychotic drugs. <u>Neuroleotic Malionant Syndrome (NMS)</u>—Potentially fatal NMS has been reported in association with administration of antipsychotic drugs, including olanzapine. See complete prescribing information for information on management of NMS. Patients requiring antipsychotic drug treatment after recovery from NMS should be carefully monitored since recurrences have been reported. <u></u>

drug discontinuation.

PIECAUTIONS: <u>Hemodynamic Effects</u>—Olarzapine may induce orthostatic hypotension associated with dizziness; tachycardia; and in some patients, syncope. Hypotension, bradycardia with/without hypotension, tachycardia, and syncope were also reported during the clinical trials with intramuscular olarzapine for injection. Incidence of syncope was 0.6%, 15/2500 with oral olarzapine in phase 2-3 trials and 0.3%, 2/722 with intramuscular olarzapine for injection in clinical trials. Three normal volunteers in phase 1 studies with intramuscular olarzapine of injection in clinical trials. Three normal volunteers in phase 1 studies with intramuscular olarzapine for injection in clinical trials. Syncope was 0.5%, 15/2500 with oral oralizatine in phase 2-3 traits and 0.3%, 2/122 with intramuscular olarazpine experienced hypotension, bradycardia, and sinus pauses of up to 6 seconds that spontaneously resolved (in 2 cases the events occurred on intramuscular olarazpine, and in 1 case, on oral olarazpine). The risk for this sequence of events may be greater in nonpsychiatric patients compared to psychiatric patients who are possibly more adapted to certain effects of psychotropic drugs. Patients should remain recumbent if drowsy or dizz atter injection with intramuscular olarazpine for injection until examination has indicated they are not experiencing postural hypotension, bradycardia, and/or hypoventilation. Olanzapine should be used with particular caution in patients with known cardiovascular disease (history of myocardia) infarction or ischemia, heart failure, or conduction abnormalities), cerebrovascular disease, and conditions which would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications) where the occurrence of syncope, or hypotension and/or bradycardia might but them at increased medical risk. Caution is necessary in patients receiving treatment with other drugs having effects that can induce hypotension, bradycardia, respiratory or ONS depression (see Drug Interactions). Concomitant administration of intramuscular olanzapine and parenteral berzodiazepine has not been studied and is not recommended. If such combination treatment is considered, cardiul evaluation of clinical status for excessive sedation and cardiorespiratory depression is recommended. <u>Seizures – During premarketing testing</u>, seizures occurred in 0.9% (22/2500) of olanzapine treated patients, regardless of causality. Use cautiously in patients with a history of seizures or with conditions that potentially lower the seizure threshold.

regardless of causality. Use cautiously in patients with a history of seizures or with conditions that potentially lower the seizure threshold. <u>Hyperprolactinemia</u>—Like other drugs that antagonize dopamine D2 receptors, olanzapine elevates prolactin levels, a modest elevation persists during chronic administration. Tissue culture experiments indicate that approximately one third of human breast cancers are prolactin dependent in vitro. However, neither clinical nor epidemiologic studies have shown an association between chronic administration of this class of drugs and tumorigenesis in humans; the available evidence is inconclusive. <u>Transaminase Elevations</u>—In placebo-controlled studies, clinically significant ALT (SGPT) elevations (s3 times the upper limit of normal) were observed in 2% (6/243) of patients exposed to blanzapine compared to no (0/115) placebo patients. None of these patients experienced jaundice. Among about 2400 patients with baseline SGPT =50 IU/L, 2% (50/2381) had asymptomatic SGPT elevations to >200 IU/L, Most were transient changes that tended to normalize while olanzapine treatment was continued. Among 2500 patients in oral olarzapine trials, about 1% (23/2500) discontinued treatment due to transaminase increases. Rare postmarketing reports of hepatitis have been received. Very rare cases of cholestatic or mixed liver injury have also been reported in the postmarketing period. Exercise caution in patients who have signs and symptoms of hepatic impairment; preexisting conditions associated with limited hepatic functional reserve; or concomitant treatment with potentially hepatotoxic drugs (see Laboratory Tests, below). <u>Potential for Coognitive and Motor impairment</u>—Somnolence was a commonly reported, dose-related adverse event in premarketing trials (olanzapine 26% vs placebo 15%). Somolence led to discontinuation in 0.4% (9/2500) of patients in the oral premarketing database. <u>Body Temperature Regulation</u>—Use, appropriate care when prescribing olanzapine for patients who will be

event in premarketing trials (planzapine 20% vs placedo 15%), somnoience led to discontinuation in 0.4% (97/500) of patients in the oral premarketing database. <u>Body Temperature Regulation</u>—Use appropriate care when prescribing olanzapine for patients who will be experiencing conditions that may contribute to an elevation in core body temperature. <u>Dysphagia</u>—Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration pneumonia is a common cause of morbidity and morbidity in patients with advanced Alcheimer's disease. Olanzapine and other antipsychotic drugs should be used cautiously in patients at risk for aspiration pneumonia. <u>Suicide</u>—The possibility of a suicide attempt is inherent in schizophrenia and in bipolar disorder, and close supervision of high-risk patients should accompany drug therapy. Prescriptions for olanzapine abuild be written for the smallest quantity of tablets consistent with good patient management. <u>Use in Patients</u> with <u>Concomitant llinesses</u>—Olanzapine a history of paratytic ileus. In 5 placebo-controlled studies in elderly patients with dementia-related psychosis (n=1184), these treatment-emergent adverse events were reported, with olanzapine at an incidence of ≥2% and significantly greater with allocaraptine risk of death compared to placebo. Olanzapine is not approved for treatment with dementia-related psychosis (n=1184), these treatment-mergent adverse events were reported, with olanzapine at an incidence of ≥2% and significantly greater with anzapine tan incidence, lettrary, increased weight, asthenia, pyrexia, pneumonia, dry mouth, visual hallicinations. Discontinuation due to adverse events was significantly greater with anzapine tan incide exprise for platents with dementia-related psychosis. If the prescriber elects to treat this patient population, vigiliance should be exercised (see BOX WARNING and WARNINGS). Because of the risk of orthostatic hypotension with olanzapine, use caution in cardiac patients (*see* Hemodynamic Ef

ZYPREXA® (Olanzapine Tablets) ZYPREXA® ZYDIS® (Olanzapine Orally Disintegrating Tablets) https://doi.org/10.1017ZYPB5245 ଆମେଧାନଃଥାର (Planzaniecfଡୀମାମହାଡ)Cambridge UnRVéରାହ AMBs

Information for Patients-See full prescribing information for information to discuss with patients taking

aboratory Tests—Periodic assessment of transaminases is recommended in patients with significant hepatic disease. Drug Interactions—Use caution when olanzapine is taken in combination with other centrally acting drugs

and alcohol. Olanzapine may enhance the effects of certain antihypertensive agents. Olanzapine may antagonize the effects of levodopa and dopamine agonists. Agents that induce CYP1A2 or glucuronyl transferase enzymes (eg. omeprazole, rifampin) may cause an increase in olanzapine clearance. Inhibitors of CYP1A2 could

the effects of levodopa and dopamine agonists. Agents that indice CYP1A2 or glucurony transferáse enzymes (eg. omeprazole, rifampin) may cause an increase in olanzapine clearance. Inhibitors of CYP1A2 could potentially inhibit olanzapine clearance. Although olanzapine is metabolized by multiple enzyme systems, induction or inhibitito of a single enzyme may appreciably after olanzapine clearance. A dosage adjustment may need to be considered with specific drugs. Activated charcoal (1 g) reduced the Cmax and AUC of oral olanzapine by about 60%. Single doses of cimetidine (800 mg) or aluminum- and magnesium-containing antacids did not affect the oral bioavailability of olanzapine. Carbamzepine (200 mg) bit) causes an approximately 50% increase in the clearance of nonzapine. Higher daily doses of carbamzepine may cause an even greater increase in olanzapine clearance. Neither ethanol (45 mg/70 kg single dose) nor warfatin (20 mg single dose) had an effect on olanzapine pharmacokinetics. Fluoxetine at 60 mg (single or multiple doses) causes a small increase in the Cmax of olanzapine and a small decrease in olanzapine; lower doses of olanzapine should be considered with avaibility between individuals, and dose motification is not routinely recommended. Fluoxoamine decreases the clearance of olanzapine; lower doses of olanzapine should be considered in patients receiving fluoxoamine concomitantly. In vitro data suggest that a clinically significant pharmacokinetics of impramine/desipramine and valproate is unlikely. Olanzapine is unlikely to cause clinically important drug interactions mediated by the enzymes CYP1A2. CYP2O9, CYP2O9, CYP2O9, corY2D6, and CYP3A. Single doses of olanzapine did not affect the pharmacokinetics of impramine/desipramine or warfarin. Multiple doses of olanzapine. Multiple doses of olanzapine did not affect the pharmacokinetics of theophylline or its metabolites. Co-administration of either diazepam or ethanol potentiated the orthostatic hypotension observed with olanzapine.

become and the provided of the polyline or its metabolites. Co-administration of intramuscular lorazepam and intramuscular olanzapine for injection added to the somnolence observed with either drug alone (see Hernodynamic Effects).
Carcinogenesis, Mutagenesis, Impairment of Fertility—The incidence of liver hemangiomas and hemangiosarcomas in female mice was significantly increased in one carcinogenicity study at 2 times the maximum human daily oral dose (MHDOD) but not in another study at 2-5 times the MHDOD (mg/m² basis). In this study there was a high incidence of early mortalities in meas in the 30/20 mg/kg/d group. The incidence of any mortalities in mais in the 30/20 mg/kg/d group. The incidence of early mortalities in mediate and end the data given olanzapine at 0.5 and 2 times the MHDOD respectively (mg/m² basis). In other studies, serum prolactin measurements of olanzapine showed elevations up to 4/oid in rats at the same doses used in the carcinogenicity studies. The relevance for human risk of the finding of prolactin mediated endocrine tumors in rodents is unknown. No evidence of mutagenic potential for olanzapine has been found.
In rats, fertility (females) and mating performance (males and females) were affected at doses 1.5-11 times the MHDOD (mg/m² basis); bierefore, olanzapine may produce a delay in ovulation.
Pregnancy Category C—There are no adequate and well-controlled studies in pregnant women. Olanzapine should be used in pregnancy only if the potential shore to the feature.
Labor and Delivery, Nursing Mothers—Parturition in rats was not affected by olanzapine with dementia-field by colanzapine is build not breast-feed.
Use in Prediatric and Ceriatric Patients—Safety and effectiveness in pediatric patients have not been established. In premarketing clinical trials in patients with schizophrenia, there was no indication of any different tolerability of lanzapine is not approved by a delay pharmeschicela substace is not approved for tratement of patients

ADVERSE REACTIONS: The following findings are based on a clinical trial database consisting of 8661 patients with approximately 4165 patient-years of exposure to oral olanzapine and 722 patients with exposure to intramuscular olanzapine for injection, including patients with schizophrenia, bipolar mania, or Alzheimer's disease Intrainuscular barizghne for injection, including patients with schizophrenia, bipolar mania, of Azhermer's disease (oral olarizghne trials) and patients with agitation associated with schizophrenia, bipolar ldisorder (manic or mixed episodes), or dementia (intramuscular olanzapine for injection trials). See the full prescribing information for details on these trials. Certain portions of the discussion below relating to dose-dependent adverse events, vital sign changes, weight dain, laboratory changes, and EGC changes are derived from studies in patients with schizophrenia and have not been duplicated for bipolar mania or agitation; however, this information is also generally applicable

details on these traits. Certain portions of the discussion below relating to dose-dependent adverse events, vital sign changes, weight gain, laboratory changes, and EGC changes are derived from studies in patients with schizophrenia and have not been duplicated for bipolar mania or agitation; however, this information is also generally applicable to bipolar mania and agitation. *Associated with Discontinuation*—Overall there was no difference in discontinuations due to adverse events in placebo-controlled oral olanzapine trials (olanzapine vs placebo: schizophrenia, 5% vs 6%; bipolar mania or agitation. *Ol* 4%; placebo 0%). Discontinuations in oral schizophrenia trials due to increases in SGPT were considered to be drug related (olanzapine 2% vs placebo 0%; see PRECAUTIONS). *Commonly Observed Adverse Events*—In 6-week, placebo-controlled, premarketing schizophrenia trials, the most common treatment-emergent adverse events associated with oral olanzapine (incidence 25% and olanzapine incidence at least twice that for placebo) were: postural hypotension, constipation, weight gain, dizzinese, personality disorder (COSTART term for nonggressive objectionable behavior), and akathisia. In 3- and 4-week placebo-controlled bipolar mania monotherapy trials, the most common treatment-emergent adverse events associated with oral olanzapine were: asthenia, dry mouth, constipation, dyspepsia, increased appetite, somolence, dizziness, and tremor. In short-term bipolar mania combination therapy trials the most common fraatment-emergent adverse events observed with olanzapine plus lithium or valeorate vere dry mouth, weight gain, incleased appetite, dizziness, back pain, constipation, dyspepsia, increased appetite; somolence, dizzines, shaft tremo in placebo-controlled trials (olanzapine for injection for agitation associated with schizophrenia or bipolar mania, sommolence was the one adverse event observed at incidence of 25% and at least twice that for placebo-controlled trials (olanzapine for injection for agitation asso

dysmenorrhea, vaginitis. Adverse Events with an Incidence ≥1% in Intramuscular Trials—The following treatment-emergent adverse events were reported at an incidence of 21% with intramuscular Janzapine for injection (2.5–10 myinjection) and at incidence greater than placebo in short-term, placebo-controlled trials in agitated patients with schizophrenia or bipolar mania: Body as a Whole—asthenia; Cardiovascular—hypotension, postural hypotension; Nervous System—somnolence, dizziness, tremor. Dose Dependency of Adverse Events in Short-Term, Placebo-Controlled Trials—Extrapyramidal Symptoms In an acute-phase controlled Clinical trial in schizophrenia, there was no significant difference in ratings scales incidence between any dose of oral olanzapine (5±2.5, 10±2.5, or 15±2.5 mg/d) and placebo for parkinsonism (Simpson-Angus Scale total score >3) or akathisia (Barnes Akathisia global score =2). In the same trial, only stathisia events (spontaneous)r eported COSTART terms akathisia and hyperkinesia) showed a statistically significantly greater adverse events incidence with the 2 higher doses of olanzapine than with placebo. The incidence of patients reporting any extrapyramidal event was significantly greater than placebo only with the

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highest dose of oral olanzapine (15±2.5 mg/d). In controlled clinical trials of intramuscular olanzapine for injection, there were no statistically significant differences from placebo in occurrence of any treatment-emergent extrapyramidal symptoms, assessed by either rating scales incidence or

Ingress tobes of value applies (152.3 fig/d). In Collidier clinical traits of infraints/clinical traitage in traiting the provided in the observe of any treatment-emergent extrapyramidal symptoms, assessed by either rating scales incidence or spontaneously reported adverse events. <u>Other Adverse Fvents</u>: Dose-relatedness of adverse events was assessed using data from this same clinical trial involving 3 fixed oral dosage ranges (5±2.5, 10±2.5, or 15±2.5 mg/d) compared with placebo. The following treatment-emergent events showed a statistically significant differences from the same clinical trial involving 3 fixed oral dosage ranges (5±2.5, 10±2.5, or 15±2.5 mg/d) compared with placebo. The events, another, termor. In a 8-week, randomized, double-blind study in patients with schizophrenia, schizophrenia, schizophrenia, schizoffective disorder comparing fixed doses of 10, 20, and 40 mg/d, statistically significant differences were seen between doses for the following: baseline to endpoint gain. 10 vs 40 mg/d: incidence of treatment-emergent prolactin elevations >24.2 ng/mL (female) or >18.77 ng/mL (male), 10 vs 40 mg/d and 20 vs 40 mg/d; fatigue, 10 vs 40 mg/d and 20 vs 40 mg/d; and diziness, 20 vs 40 mg/d. In discipance or land anazpine for injection was associated with bradycardia, hypotension, and tackycardia in clinical trials (see PRECAUTIONS). *Wital Sign Changes*—Oral olanzapine for injection was associated with bradycardia, hypotension, and tackycardia in clinical trials (see PRECAUTIONS). *Wital Sign Changes*—Oral olanzapine patients gaine ornpared to 0.8% of placeho patients (average 4.4% gain) compared to 0.8% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compared to 0.3% of placeho patients (average 4.4% gain) compa

2.2.4 respectively, in these solution a mean baseline of 203 mg/dL, which was significantly different compared to placebo-treated patients (N=1415) with a mean decrease of 4.6 mg/dL from a mean baseline of 203 mg/dL.
 EG Changes—Analyses of pooled placebo-controlled trials revealed no statistically significant olanzapine/placebo differences in incidence of potentially important changes in EGG parameters, including QT, OTc, and PR intervals. Olanzapine/placebo differences in incidence of potentially important changes in EGG parameters, including QT, OTc, and PR intervals. Olanzapine **JTrials**—The following treatment-emergent events were reported with oral olanzapine placebo patients.
 Other Adverse Events Observed During Clinical Trials—The following treatment-emergent events were reported with oral olanzapine tar multiple doses >1 mg/d in clinical trials (8661 patients, 1456 patient-years of exposure). This ist may not include events previously listed elsewhere in labeling, those events for which a drug cause was remote, those terms which were as opereral as to be uninformative, and those events reported only once or twice which did not have a substantial probability of being acutely life-threatening. *Frequent:* acutal pain, flu syntome, Infrequent: abarted cause was remote, those terms which were so general as to be uninformative, and those events reported only once or twice which did not have a substantial usic occurred in 1/100 to 1/1000 patients, infrequent: abarted patients, and events, page terms, heart rates theorem trage, migriant, neak ng/dy reliation, waschildation, which are dema, which and in the statemeter, page terms, heart rates theorem trage, migrianto, masol latation, wentricular extrasystoles; *Rare:* attertits, heart failure, patient, and yoer refect, sudden deation, carreitage, migriants, heart states of the states of theartes, theoremose, and theartes, and the states of the states

Introducts enarged, vaginal nenormage , have: audministria, oreast enargement, instituts, original: (*Adjusted for gender.) The following treatment-emergent events were reported with intramuscular olanzapine for injection at one or more doses a≥2 mg/nijection in clinical trials (722 patients). This list may not include events previously listed elsewhere in labeling, those events for which a drug cause was remote, those terms which were so general as to be uninformative, and those events reported only once or twice which did not have a substantial probability of being acutely life-threatening. Body as a Whole—Frequent: injection site pain; *Intrequent*: addominal pain, fever. Cardiovascular—infrequent: AV block, heart block, syncope. Digestive—infrequent: creatine phosphokinase increased, dehydration, hyperkalemia. Musculoskeletal—infrequent: witching. Mervaus System—infrequent: abnormal gait, akthisia, articulation impairment, contusion, emotional lability. Skin and Appendages—infrequent: sweating. Postintroduction Reports—Reported since market introduction and temporally (not necessarily causality) related to olanzapine therapy: allergic reaction (e.g., anaphylactoid reaction, angioedema, purulus or uticaria), diabetic coma, jaundice, neutropenia, pancreatitis, priapism, rhabdomyolysis, and venous thromboembolic events (including pulmonary embolism and deep venous thromboesis). Random cholesterol levels of ≥240 mg/dL and random triglyceride levels of ≥1000 mg/dL have been reported. DRUG ABUSE AND DEFENDENCE: Olanzapine is not a controlled substance.

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PRIMARY PSYCHIATRY

BRIEF SUMMARY. See package insert for full prescribing information.

Increased Mortality in Elderly Patients with Dementia-Related Psychosis: Elderly patients with dementia-related psychosis treated with a typical antipsycholic drugs are at an increased risk of death compared to placebo. Analyses of seventeen placebo controlled trials (modal duration of 10 weeks) in these patients revealed a risk of death in the drug-treated patients of between 1.6 to 1.7 times that seen in placebo-treated patients. Over the course of a typical 10 week controlled trial, the rate of death in drug-treated, maint of the dentia supeared to be either cardiovascular (e.g., heart failure, sudden death) or infectious (e.g., pneumonia) in nature. GEODON (ziprasidone) is not approved for the treatment of patients with Dementia-Related Psychosis.

NDICATIONS--GEODON Capsules is indicated for the treatment of schizophrenia and acute manic or mixed episodes associated with

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Control acute agitation with GEODON for **Injection** (ziprasidone mesylate)

In schizophrenia... Rapid control* with low EPS¹⁻⁴

- Low incidence of movement disorders¹⁻⁴
- Smooth transition, with continued improvement, from IM to oral therapy^{3,4}
- May be used concomitantly with benzodiazepines^{2,3,5}

* In 2 pivotal studies vs control, significance was achieved at the 2-hour primary end point (10 mg study) and at the 4-hour primary end point (20 mg study).



GEODON for Injection is indicated for the treatment of acute agitation in schizophrenic patients for whom treatment with GEODON is appropriate and who need intramuscular antipsychotic medication for rapid control of the agitation.

Elderly patients with dementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk of death compared to placebo. GEODON is not approved for the treatment of patients with dementia-related psychosis.

GEODON is contraindicated in patients with a known history of QT prolongation, recent acute myocardial infarction, or uncompensated heart failure, and should not be used with other QT-prolonging drugs. GEODON has a greater capacity to prolong the QT_c interval than several antipsychotics. In some drugs, QT prolongation has been associated with torsade de pointes, a potentially fatal arrhythmia. In many cases this would lead to the conclusion that other drugs should be tried first.

As with all antipsychotic medications, a rare and potentially fatal condition known as neuroleptic malignant syndrome (NMS) has been reported with GEODON. NMS can cause hyperpyrexia, muscle rigidity, diaphoresis, tachycardia, irregular pulse or blood pressure, cardiac dysrhythmia, and altered mental status. If signs and symptoms appear, immediate discontinuation, treatment, and monitoring are recommended. Prescribing should be consistent with the need to minimize tardive dyskinesia (TD), a potentially irreversible dose- and duration-dependent syndrome. If signs and symptoms appear, discontinuation should be considered since TD may remit partially or completely.

Hyperglycemia-related adverse events, sometimes serious, have been reported in patients treated with atypical antipsychotics. There have been few reports of hyperglycemia or diabetes in patients treated with GEODON, and it is not known if GEODON is associated with these events. Patients treated with an atypical antipsychotic should be monitored for symptoms of hyperglycemia.

Precautions include the risk of rash, orthostatic hypotension, and seizures. In fixed-dose, pivotal studies, the most commonly observed adverse events associated with the use of GEODON for Injection (incidence \geq 5%) and observed at a rate in the higher GEODON dose groups (10 mg, 20 mg) of at least twice that of the lowest GEODON dose group (2 mg control) were somnolence (20%), headache (13%), and nausea (12%).

Please see brief summary of prescribing information on adjacent page.

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The International Journal of Neuropsychiatric Medicine

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Paul E. Keck, Jr, MD, Roger S. McIntyre, MD, FRCPS, and Richard C. Shelton, MD

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