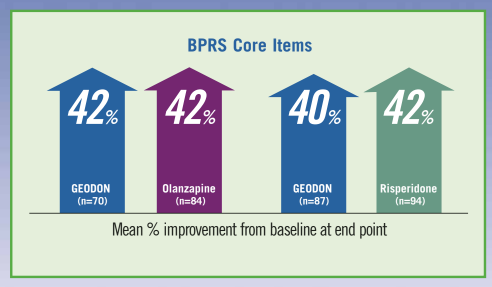
Treat schizophrenia

COMPARABLE EFFICACY

Consistent results in head-to-head studies1-3



A 6-week, double-blind, randomized study of GEODON vs olanzapine and an 8-week, double-blind, randomized study of GEODON vs risperidone.

- BPRS core items include hallucinatory behavior, unusual thought content, conceptual disorganization, and suspiciousness
- Comparable efficacy was maintained in double-blind extension studies
 - —up to 1 year vs risperidone¹
 - —up to 6 months vs olanzapine⁴

GEODON is indicated for the treatment of schizophrenia and of acute manic or mixed episodes associated with bipolar disorder.

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.

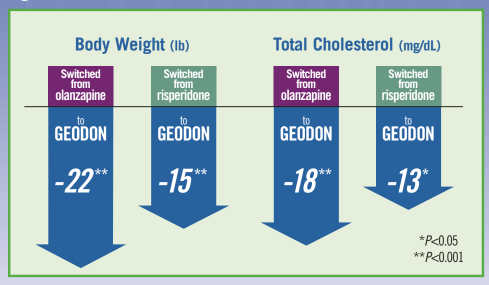
Hyperglycemia-related adverse events, sometimes serious, have been reported in patients treated with atypical antipsychotics. There have been few reports of hyperglycemia or diabetes in patients treated with GEODON, and it is not known if GEODON is associated with these events. Patients treated with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

In short-term schizophrenia trials, 10% of GEODON-treated patients experienced a weight gain of $\geq 7\%$ of body weight vs 4% for placebo. In the same short-term trials, the most common adverse events were somnolence (14%) and respiratory tract infection (8%).

with the body in mind

WITHOUT COMPROMISING METABOLIC PARAMETERS

Significant results in switch studies^{1,5}



Two 1-year open-label extensions of 6-week, open-label switch studies in patients suboptimally controlled due to partial response or poor tolerability.

■ Patients switching to GEODON from olanzapine and risperidone also experienced reductions in triglycerides⁵

In the acute head-to-head studies...

- In the GEODON vs olanzapine study, olanzapine significantly increased body weight (8 lb vs 2 lb for GEODON, *P*<0.0001)^{1,2}
- In the GEODON vs risperidone study, risperidone increased body weight (2 lb vs 0 lb for GEODON, *P*<0.01)¹,³





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 (e.g., heart failure, sudden teath) or intectious (e.g., pneumonia) in nature. GEODON (ziprasidone) is not approved for the treatment of natients with Dementia-Related Psychosis.

INDICATIONS—GEODON Capsules is indicated for the treatment of schizophrenia and acute manic or mixed episodes associated with bipolar disorder with or without psychotic features. GEODON® (ziprasidone mesylate) for Injection is indicated for acute agitation in schizophrenic analients.

CONTRAINDICATIONS—QT Prolongation: Decause of GEODON's dose-related prolongation of the QT interval and the known association of fatal arrhythmias with QT prolongation by some other drugs, GEODON is contraindicated in patients with a known history of QT prolongation (including congenital long QT syndrome), with recent acute myocardial infarction, or with uncompensated heart failure (see WARNINGS). Pharmacokinetic/pharmacodynamic studies between GEODON and other drugs that prolong the QT interval have not been performed. An additive effect of GEODON and other drugs that prolong the QT interval cannot be excluded. Therefore, GEODON should not be given with dofetilide, sotaloi, quinidine, other Class Ia and III anti-arrhythmics, mesoridazine, thioridazine, chlorpromazine, droperidol, pimozide, sparfloxacin, gatifloxacin, moxifloxacin, moxifloxacin, halofantrine, mefloquine, pentamidine, arsenic trioxide, levomethadyl acetate, dolasetror mesylate, probucol, or tacrolimus. GEODON is also contraindicated with drugs that have demonstrated QT prolongation as one of their nesynate, produced, in accommus, seven but effect described in the full prescribing information as a contraindication or a boxed or bolded warning (see WARNINGS). GEODON is contraindicated in individuals with a known hypersensitivity to the product. WARNINGS—Increased Mortality in Editory Patients with Dementia-Related Psychosis: Editory patients with thementia-related psychosis treated with atypical antipsychotic drugs are at an increased risk of death compared to placebo. GEODON (ziprasidone) is not approved for the treatment of patients with dementia-related psychosis (see Boxed Warning). 2T Prolongation and Risk of Sudden Death: GEODON us should be avoided in combination with other drugs that are known to prolong the QT, interval. Additionally, clinicians should be alert to the identification of other drugs that have been consistently observed to prolong the QT, interval. Additionally, clinicians should be alert to the identification of other drugs that have been consistently observed to prolong the QT, interval. Such drugs should not be prescribed with GEODON. A study directly comparing the QTQT_prolonging effect of GEODON with several other drugs effective in treatment of schizophrenia was conducted in patient volunteers. The mean increase in QT, from baseline for GEODON and rugs effective in treatment of schizophrenia was conducted in patient volunteers. The mean increase in QT, from baseline for GEODON and haloperidol), but was approximately 4 hasse less than the prolongation hoserved for thirdrazine. In this study, the effect of GEODON with certification of the properties of the pharmacodynamic effects and have this effect described in the full prescribing information as a contraindication or a boxed or bolded warning Intereationship of ut protongation to torsade ae pointes is clearest for larger increases (20 misee and greater) but it is possible that smaller OT/CT, protongations may also increase risk, or increase it in susceptible individuals, such as those since and preater but it is possible that small protonging effect of intermuscular protonging effect of intramuscular GEODON, with intramuscular haloperidol (3 as a control, was conducted in patient volunteers. In the trial, EGS were obtained at the time of maximum plasma concentration following two injections of EEDDON (20 mg then 30 mg) or haloperidol (7.5 mg then 10 mg) given four hours apart. Note that a 30 mg dose of intramuscular GEODON is 50% higher than the recommended therapeutic dose. The mean change in OT_c from baseline was calculated for each drug using a sample-based correction that removes the effect of heart rate on the OT interval. The mean increase in OT_c from baseline for GEODON was 4.6 mise following the first injection and 12.8 mise following the second injection. The mean increase in OT_c from baseline for haloperidol was 6.0 misec following the second injection in this study, no patient had a OT_c interval exceeding 500 misec. As with other antipsychotic drugs and placebo, sudden unexplained details have been reported in patients taking GEODON at recommended doses. The premarketing experience for GEODON did not reveal an excess of mortality for GEODON compared to other antipsychotic drugs and placebo, sudden unexplained details have been reported in patients taking GEODON. At recommended doses. The premarketing experience for GEODON did not reveal an excess of mortality for GEODON compared to other antipsychotic drugs and placebo, but the extent of exposure was limited, especially for the drugs used as active controls and placebo. Nevertheless, GEODON's targer prolongation of OT_c length compared to several other antipsychotic drugs are set to exceed the possibility heads to be considered in deciding among alternative drug products. Certain patients being considered for GEODON treatment who are at risk for significant electrolyte disturbances, hypokalemia in particular, have baseline serum potassium and magnesium measurements. Hypokalemia, and/or hypomagnessemia) may inceste the risk of QT prolongation and arrhythmia. Hypokalemia may result from diuretic therapy, diarrhea, and other causes. Patients with low serum potassium and/or magnesium should be repleted with those electrolytes before proceeding with treatment. It is essential to periodically monitor serum electrolytes in patients for whom diuretic therapy is infroduced during GEODON treatment. Persistently prolonged QT_c intervals may also increase the risk of further prolongation and arrhythmia, but it is not clear that routine screening ECG measures are effective in delection such patients. Rather, GEODON should be avoided in patients with histories of significant cardiovalar illness, eg, QT prolongation, recent acute myocardial infarction, uncompensated heart failure, or cardiac arrhythmia. GEODON should be avoided in patients with histories of significant cardiovalar illness, eg, QT prolongation, recent acute myocardial infarction, uncompensated heart failure, or cardiac arrhythmia. GEODON should be avoided in patients with histories of significant cardiovalar illness, eg, QT prolongation, recent acute myocardial infarction, uncompensated heart failure, or cardiac arrhythmia. GEODON should be avoided in patients who are found to have persistent QT, measurements > 500 msec. Neuroleptic Malignant Syndrome (MMS): A potentially fatale spymptom complex sometimes referred to as Neuroleptic Malignant Syndrome (MMS): As been reported in association with administration of antipsychotic drugs. The management of MMS should include: (1) immediate discontinuation of antipsychotic drugs. The management of MMS should include: (1) immediate discontinuation of antipsychotic drugs. The management of MMS should include: (1) immediate discontinuation of antipsychotic drugs. The management of MMS is should auministration of an applycrotic drugs. The intelligenter into which stoud include: (1) minied as described become an other drugs not sesential to concurrent therapy. (2) intensive symptomatic treatment and medical monitoring; and (3) treatment of any concomitant serious medical problems for which specific treatments are available. If a patient requires antipsychotic drug treatment after recovery from NMS, the potential reintroduction of drug therapy should be carefully considered. The patient should be carefully monitored, since recurrences of NMS have been reported. Tardive Dyskinesia (TD): A syndrome of potentially inversible, involuntary, dyskinetic movements may develop in patients undergoing treatment with antipsychotic drugs. Although the prevalence of TD appears to be higher among the elderly, especially deldry women, it is impossible to rely upon prevalence estimates to predict, at the inception of antipsychotic treatment, which patients are likely to develop TD. If signs and symptoms of TD appear in a patient on GEODON, drug discontinuation should treatment, which patients are likely to develop TD. If signs and symptoms of TD appear in a patient on GEODON, drug discontinuation should be considered. *Hyperglycemia and Diabetes Mellitus: Hyperglycemia-related adverse events, sometimes serious, have been reprorted 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 by ymptoms of hyperglycemia. PreEQUITIONS — *General: Rash: In premarketing trials, about 5% of GEODON patients developed rash and/or urbicaria, with discontinuation of treatment in about one-sixth of these cases. The occurrence of rash was dose related, although the finding might also be explained by longer exposure in higher-dose patients. Several patients with rash had signs and symptoms of associated systemic illness, e.g., elevated MVBCs. Most patients improved promptly upon treatment with antibiastamines or steroids and/or upon discontinuation of GEODON, and all patients were reported to recover completely. Upon appearance of rash for which an alternative etiology cannot be identified, GEODON should be discontinued. *Orthostatic Hypotension: GEODON in prior or orthostatic hypotension is orthostatic hypotension prior orthostatic hypotension prior orthostatic hypotension is orthostatic hypotension or orthostatic hypotension is orthostatic hypotension prior orthostat with dizziness, tachycardia, and, in some patients, syncope, esspecially during the initial dose-titration period, probably reflecting its c₁-adrenergic antagonist properties. Syncope was reported in 0.6% of GEODON patients. GEODON should be used with particular caution in patients with known cardiovascular disease (history of myocardial infarction or ischemic heart disease, heart failure or conduction abnormalities), cerebrovascular disease or conditions that would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications). Seizures: In clinical trials, seizures occurred in 0.4% of GEODON patients. There were confounding factors that may have contributed to seizures in many of these cases. As with other antipsychotic drugs, GEODON should be used cautiously in patients with a history of seizures or with conditions that potentially lower the seizure threshold, e.g., Atzheimer's dementia. Conditions that lower the seizure threshold may be more prevalent in a population of 65 years or older. <u>Dysphagia:</u> Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration pneumonia is a common cause of motificity and mortality elderly patients, in particular those with advanced Atcheimer's dementia, and GEODOM and other antipsychotic drugs should be used cautiously in patients at risk for aspiration pneumonia. (See also Boxed WARNING, WARNINGS: Increased Mortality in Elderly Patients with Dementia-Related Psychosis, Hyperprotectinemia: As with other drugs that antagonize dopamine D₂ receptors, GEODON elevates protactin levels in humans. Tissue culture experiments indicate that approximately one third of human breast cancers are protactin dependent in vitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancer. In Mind of a reactive to personal importance in the prescription or insecurings some impact with a presume whether the contract shudies not epidemiologic shudies conducted to date have shown an association between chronic administration of this class of drugs and tumorigenesis in humans; the available evidence is considered too limited to be conclusive at this time. Potential for Cognitive and Motor Impairment. Somnolence was a commonly reported adverse event in GEODON patients. In the 4- and 6-week placebo-controlled trials, somnolence was reported in 14% of GEODON patients vs 7% of placebo patients. Somnolence led to discontinuation in 0.3% of patients in short-term clinical trials. Since GEODON has the potential to impair judgment, thinking, or motor skills, patients should be cautioned about performing activities requiring mental alertness, such as operating a motor vehicle (including automobiles) or operating hazardous machinery until they are reasonably certain that GEODON therapy does not affect them adversely. <u>Priapism: One case of priapism</u> was reported in the premarketing database. <u>Body Temperature Regulation:</u> Although not reported with GEODON in premarketing trials, disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents. <u>Suicide:</u> The possibility of a suicide attempt is inherent in psychotic illness and close supervision of high-risk patients should accompany drug therapy. GEODON prescriptions should be written for the smallest quantity of capsules consistent with good patient management to reduce overdose risk. Use in Patients with Concomitant Illness: Clinical experience with GEODON in patients with certain concomitant systemic illnesses is limited. GEODON has not been evaluated or used to any appreciable extent in patients with a recent history of myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from premarketing clinical studies. Because of the risk of OT_c prolongation and orthostatic hypotension with GEODON, caution should be observed in cardiac patients (see *QT Prolongation and Risk of Sudden Death* in WARNINGS and Orthostatic Hypotension in PRECAUTIONS). Information for Patients: To ensure safe and effective use of GEODON, the

information and instructions in the Patient Information Sectionshould be discussed with patients. Laboratory Tests: Patients being considered for GEODON treatment who are at risk of significant electrolyte disturbances should have baseline serum potassium and magnesium measurements. Low serum potassium and magnesium should be repleted before treatment. Patients who are started on diuretics during GEODON therapy need periodic monitoring of serum potassium and magnesium. Discontinue GEODON in patients who are started online to have persistent OT_c measurements >500 msec (see **WARNINGS**). *Drug Interactions*: (1) GEODON should not be used with any drug that prolongs persistent of, measurements Sout mise, see Warkinson, brighted in the CT interval (2) Given the primary CNS effects of GEODON, caution should be used when it is taken in combination with other centrally acting drugs. (3) Because of its potential for inducing hypotension, GEODON may enhance the effects of certain antihypertensive agents. (4) GEODON may antagonize the effects of levodops and doparnine agonists. Effect of Other Drugs on GEODON. Carbamazepine, 200 mg off or 2 days, resulted in a decrease of approximately 35% in the AUC of GEODON. Retoconazole, a potent inhibitor of CYP3A4, 400 mg off or 5 days, increased the AUC and C_{max} of GEODON by about 35% 40%. Cimetidine, 800 mg of for 2 days, did not affect GEODON pharmacokinetics. Coadministration of 30 mL of Meakov did not affect GEODON pharmacokinetics. Population pharmacokinetics are applied to the production of the pharmacokinetic analysis of the pharmacokinetics of the pharmacokinetics of the pharmacokinetics are applied to the pharmacokinetics of the pharmacokinetics are applied to the pharmacokinetics of the pharmacokinetic analysis of the pharmacokinetics of the pharmacokinetics and the pharmacokinetics of the pharmacokinetic analysis of the pharmacokinetic analys or Schizophrenic patients in controlled clinical trials has not revealed rey clinically significant pharmacokinetic interactions with benztropine, propranolol, or lorazepam. Effect of GEODON on Other Drugs; In vitro studies revealed little potential for GEODON to interfere with the metabolism of drugs cleared primarily by CYP142, CYP205, CYP205, CYP205, GYP205, and CYP344, and little potential for GEODON to interfere with the metabolism of drugs cleared primarily by CYP142, CYP205, There was no statistically significant change in the urinary dextromethorphan/dextrorphan ratio. Carcinogenesis, Mulagenesis, Impairment of Fertility: Lifetime carcinogenicity studies were conducted with GEODON in Long Evans rats and CD-1 mice. In male mice, there was no increase in incidence of tumors relative to controls. In female mice there were dose-related increases in the incidences of pitulary gland adenoma and carcinoma, and mammary gland adenocarcinoma at all doses tested. Increases in retirection were observed in a 1-month dietary study in female, but not male, mice. 6EODON had no effect on serum protactin in a 5-week dietary study at the doses that were used in the carcinogenicity study. The relevance for human risk of the findings of protactin-mediated endocrine tumors in rodents is unknown (see https://perpotactinemia). Mutagenesis: There was a reproducible mutagenic response in the Ames assay in one strain of S. *pphimumium in the absence of metabolic activation. Positive results were obtained in both the in vitro mammalian cell gene mutation assay and the in vitro chromosomal aberration assay in human hymphocytes. Impairment of Fertility, GEODON increased time to copulation in Sprague-Dawley rats in two fertility and early embryonic development studies at doses of 10 to 160 mg/kg/day (0.5 to \$10 to \$10 to \$10 mg/kg/day (0.5 to \$10 to \$10 mg/kg/day (0.5 to \$10 to \$10 to \$10 mg/kg/day (0.5 to \$10 to 8 times the MRHD of 200 mg/day on a mg/m² basis). Fertility rate was reduced at 160 mg/kg/day (8 times the MRHD on a mg/m² basis). There was no effect on fertility at 40 mg/kg/day (2 times the MRHD on a mg/m² basis). There was no effect on fertility at 40 mg/kg/day (2 times the MRHD on a mg/m² basis). The fertility of female rats was reduced. **Pregnancy Category C:** There are no adequate and well-controlled studies in pregnant women. GEODON should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Labor and Delivery: The effect of GEODON on labor and delivery in humans is unknown. Nursing Mothers: It is not known whether, and if so in what amount, GEODON or its metabolites are excreted in human fit is recommended that women receiving GEODON should not breast feed. Pediatric Use: The safety and effectiveness of GEODON in pediatric patients have not been established. Geriatric Use: Of the approximately 4500 patients treated with GEODON in clinical studies, 2.4% (109) were 65 years of age or over. In general, there was no indication of any different tolerability for GEODON or freduced clearance of GEODON in the elderly compared to younger adults. Nevertheless, the presence of multiple factors that might increase the pharmacodynamic response to GEODON, or cause poorer tolerance or orthostasis, should lead to consideration of a lower starting dose, pharmacodynamic resporse to GEODON, or cause poorer tolerance or orthostasis, should lead to consideration of a lower starting dose, sower titeration, and careful monitoring during the initial dosing period for some elderly patients. AUVERSE REACTIONS—Aures Findings Observed in Short-term, Placebo-Controlled Trials: The following findings are based on the short-term placebo-controlled premarketing trials for schizophrenia (a pool of two 6-week, and two 4-week fixed-dose trials) and bipolar mania (a pool of two 3-week fixeible-dose trials) in which GEODON was administered in doses ranging from 10 to 200 mg/day. Adverse Events Associated with Discontinuation: Schizophrenia: Approximately 4.1% (29702) of GEODON reated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 2.6% (6273) on placebo. The most common event associated with orthou was rash, including 7 dropouts for rash among GEODON patients (1%) compared to no placebo patients (see PRECAUTIONS). Bipolar Mania: Approximately 6.5% (18279) of GEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 3.7% (5/136) on placebo. The most common events associated with dropout in the GEODON-treated patients were akathisia, anxiety, depression, dizziness, dystonia, rash and vorniting, with 2 dropouts for each of these events among GEODON patients (1%) compared to the placebo Dadent adae to no lacebo patients for the remaining three events. patients (1%) compared to one placebo patient each for dystonia and rash (1%) and no placebo patients for the remaining a diverse events.

**Adverse Events at an Incidence 25% and at Least Twice the Rate of Placebo. The most commonly observed adverse events associated with 6EODON in schizophrenia trials were sonnolence (14%) and respiratory tract infection (8%). The most commonly observed adverse events associated with the use of GEODON in bipolar mania trials were sonnolence (31%), extrapyramidal symptoms (31%), dizziness events associated with the use of GEODON in bipolar main at irals were somnolence (31%), extrapyramidal symptoms (31%), duziness (16%), adurations (10%), abnormal vision (6%), adbraich (10%), abnormal vision (6%), adbraich (36%), and womiting (5%). The following list enumerates the treath-emergent adverse events that occurred uring acute therapy, including only those events that occurred in 2% of GEODON patients and at a greater incidence than in placebo. Schizophrenia: Body as a Whole—asthenia, accidental injury, chest pain. Cardiovascular—tachycardio, <u>Diegstive—nausea</u>, constipation, dyspepsia, diarrhea, dry mouth, anorevia. <u>Nervous—extrapyramidal symptoms</u>, somnolence, akathisia, dizziness. <u>Respiratory—respiratory tract infection</u>, thinitis, cough increased. <u>Skin and Appendages—rash</u>, flung dermatitis. <u>Special Senses—abnormal vision</u>, Bipolar Mania: <u>Body as a Whole—headache</u>, asthenia, accidental injury, <u>Cardiovascular—myyerlension</u>. <u>Digestive—nausea</u>, diarrhea, dry mouth, vomiting, increased salivation, tongue edema, dysphagia. <u>Musculoskeletal—myalgia. Nervous—somnolence</u>, extrapyramidal symptoms, disziness, akathisia, anxiety, hypesthesia, speech disorder. <u>Respiratory—pharyngits</u>, dyspnea. <u>Skin and Appendages—tungal dematitis. Special Senses—abnormal vision. Digestive—nausea</u>, diarrhea, and paperant relation of adverse event to dose for the following: asthenia, postural hypotension, anorexia, driviness, dysfonia bunertonia somnolence tempor chimitis resh, and ahonormal vision. mouth, increased salivation, arthraligia, anxiety, dizziness, dystonia, hypertonia, somnolence, tremor, rhinitis, rash, and abnormal vision.

Extrapyramidal Symptoms (EPS): The incidence of reported EPS for GEODON patients in the short-term, placebo-controlled schizophrenia trials was 14% vs 8% for placebo. Objectively collected data from those trials on the Simpson-Angus Rating Scale and the Barnes Akathisia trials was 14% vs 8% for placebo. Objectively collected data from those trials of the elimptonia phase and the parties are stated and the parties and the sames Aratively collected data from those trials of the elimptonia phase (ECDON) and placebo. Wila Sign Changes: (ECDON) is associated with orthostatic hypotension (see PRECAUTIONS). Weight Gain: In short-term schizophrenia trials, the proportions of patients meeting a weight gain criterion of 27% of body weight were compared, revealing a statistically significantly greater incidence of weight gain for GEDON patients (10%). A median weight gain of 0.5 kg was observed in GEDON patients vs 0.0 kg in placebo patients. Weight gain was reported as an adverse event in 0.4% of both GEDON and placebo patients. During long-term therapy with GEDON activorsity of patients at baseline on the basis of body mass index (RMI) showed the greatest mean weight gain and the light patient of the patients are considered as an adverse event in 0.4% of both GEDON and placebo patients. During long-term therapy with GEDON and placebo patients. During long-term therapy with GEDON and placebo patients. During long-term therapy with GEDON and placebo patients. patients. There was a mean weight gain of 1.4 kg for patients with a "low" baseline BMI, 0.0 kg for patients with a "hom?" BMI, and a 1.3 kg mean weight loss for patients with a "high" BMI. **EGG Changes**: GEODON is associated with an increase in the 0.1_c interval (see **WARNINGS**). In schizophrenia trials, GEODON was associated with a mean increase in heart rate of 1.4 beats per minute compared to a 0.2beats per minute decrease among placebo patients. *Other Adverse Events Observed During the Premarketing Evaluation of GEODON:*Frequent adverse events are those occurring in at least 1/100 patients, infrequent adverse events are those occurring in 1/100 to 1/1000 patients; rare events are those occurring in fewer than 1/1000 patients. Schizophrenia: <u>Body as a Whole</u>—*Frequent:* abdominal pain, flu syndrome, fever, accidental fall, face edema, chills, photosensitivity reaction, flank pain, hypothermia, motor vehicle accident. Cardiovascular System—Frequent: tachycardia, hypertension, postural hypotension; Infrequent: bradycardia, angina pectoris, atrial fibrillation; Rare: first-degree AV block, bundle branch block, phlebitis, pulmonary embolus, cardiomegaly, cerebral infarct, cerebrovascular accident, deep tegree voluce, dunie traint notes, imientis, pluminary entoutos, catoninegari, creatian interio, ceretata interior, teretata hemorrhage, intromotophiebitis, myocarditis, thrombophiebitis, <u>Digestive System — Frequenit anorexia</u>, vomiting, *Infrequenit rectata* hemorrhage, dysphagia, tongue edema; *Rare*; gum hemorrhage, jaundice, fecal impaction, gamma glutamy transpeptidase increased, hematemest, colestatic jaundice, hepatitis, hepatomegaly, leukoplakis of mouth, fatly liver deposit, melena <u>Endocrine</u>——*Rare*: hypothyroidism, hyperthyroidism, thyroiditis. <u>Hemic and Lymphatic System — Infrequent</u>: anemia, ecchymosis, leukocytosis, leukopenia, eosinophilia, lymphadenopathy, *Rare*: thrombocytopenia, hypochromic anemia, lymphocytosis, monocytosis, basophilia, lymphadema, polycythemia, thrombocythemia, Metabolic and Nutritional Disorders — *Infrequent*: thirst, transaminase increased, peripheral edema, polycythemia, creatine phosphokinase increased, alkaline phosphatase increased, hypercholesteremia, dehydration, lactic dehydrogenase increased. adamin prospriorinas incleased, anamine prospriadase incleased, hyperlipenia, hypocholesteremia, hyperlalemia, hypocholesteremia, hypocholesteremi oupopa, incoordination, neuropatiny, interupent: parayisis, Faze: myoconus, nystagmus, torticosis, circumora parasinesa, opisionotoris, reflexes increased, trismus. Reportatory <u>System - Frequent dyspase</u>, *Infraquent*, proaumonia, epistasis, Faze: hemopytosis, larynipismus. Skin and <u>Appendages — Infraquent</u> maculopapular rash, urticaria, alopecia, eczema, exfoliative dermatitis, contact dermatitis, vesiculobullous rash. <u>Special Senses — Frequent</u> trugal elematitis, *Infraquent* comjunctivis, dry eyes, tinnitus, blepharitis, cataract, photophobia, *Paze* eye hemorrhage, visual field defect, keratitis, keratoconjunctivitis. <u>Urogenital System — Infraquent</u> impotence, abnormal ejaculation, amenorrhage, hematuria, menorrhage, female detaction, polyuria, urinary retention, metrorrhagia, male sexual dysfunction, anorgasmia, glycocounia; *Paze* eyencomastia, vaginal hemorrhage, nocturia, oliginari, female sexual dysfunction, theremorrhage. **Adverse Finding Observed in Trials of Inframuscular GEODON**: In these studies, the most commonly observed adverse events associated with he use of intramuscular GEODON is the biober dose grower blace this contramuscular GEODON. with the use of intramuscular GEODON (25%) and observed at a rate on intramuscular GEODON (in the higher dose groups) at least twice that of the lowest intramuscular GEODON group were headache (13%), nausea (12%), and somnolence (20%). Adverse Events at an Incidence >1% in Short-Term Fixed-Dose Intramuscular Trials: The following list enumerates the treatment-emergent adverse events Incidence 51% in Short-Term Fixet-Dose Intramuscular Inals: The following list enumerates the freatment-emergent adverse events that occurred in a 1% of 6EODON patients (in the higher dose groups) and at least twice that of the lowest inframuscular poble intermous many patients (in the patient state). By the patient of the patient state of the patients of the pati mg), the only symptoms reported were minimal sedation, slurring of speech, and transitory hypertension (BP 200/75)

References: 1. Data on file. Pfizer Inc., New York, NY. 2. Simpson GM, Glick ID, Weiden PJ, Romano SJ, Siu CO, Randomized, controlled, double-blind multicenter comparison of the efficacy and tolerability of ziprasidone and olarazapine in acutely ill inpatients with schizophrenia or schizoaffective disorder. Am J Psychiatry, 2004;161:1837-1847. 3. Addington DEN, Pantelis C, Dineen M, Benattia I, Romano SJ. Efficacy and tolerability of ziprasidone versus risperidone in patients with acute exacerbation of schizophrenia or schizoaffective disorder: an 8-week, double-blind, multicenter trial. J Clin Psychiatry, 2004;65:1624-1633. 4. Simpson GM, Weiden P, Pigott T, Murray S, Siu CO, Romano SJ. Six-month, blinded, multicenter continuation study of ziprasidone versus olanzapine in schizophrenia. Am J Psychiatry, 2005;162:1535-1538. 5. Weiden PJ, Loebel A, Yang R, Lebovitz H. Course of weight & metabolic benefits 1 year after switching to ziprasidone. Presented at: American Psychiatric Association Annual Meeting: May 1-6, 2004; New York, NY.

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The 2nd Annual Advances in Psychiatry: A Guide to Clinical Practice April 27 & 28, 2007

at the

Grand Hyatt New York New York, New York

Program Description & Objectives

This conference is designed for psychiatrists in clinical practice and other mental health professionals involved in the treatment of psychiatric disorders in adults. The faculty of this conference will include members of the Department of Psychiatry at Columbia University College of Physicians and Surgeons and the New York State Psychiatric Institute. Faculty involved in research and academic programs will address common clinical dilemmas with practical solutions informed by available evidence. Course participants will acquire knowledge about the diagnosis and treatment of the major psychiatric illnesses in adults, with a focus on the integration of recent research findings in clinical decision-making. At the conclusion of the program they will be better able to use research findings to manage clinical situations and to evaluate new data as it emerges.

> Course Directors: Frederic Kass, M.D., Jeffrey A. Lieberman, M.D., Ronald O. Rieder, M.D. & Mary S. Sciutto, M.D.

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Important Safety Information for ZYPREXA® (olanzapine)

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 drugtreated 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 drugtreated 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 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 elderly 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.

Tardive dyskinesia (TD)—As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of TD. If its signs and symptoms appear, discontinuation should be considered.

Orthostatic hypotension—In premarketing schizophrenia trials, some patients taking ZYPREXA may have experienced orthostatic hypotension associated with dizziness, tachycardia, and, in some cases, syncope (15/2500, 0.6%).

Seizures—Occurred infrequently in premarketing clinical trials (22/2500, 0.9%). ZYPREXA should be used cautiously in patients with a history of seizures or with conditions that lower the seizure threshold.

Effect on prolactin—Modest elevations of prolactin were seen with ZYPREXA in acute-phase schizophrenia trials (incidence 34% vs 13% with placebo), although mean changes from baseline to endpoint were not statistically significantly different between olanzapine and placebo. Some patients may have persisting modest prolactin elevations.

Transient, asymptomatic elevations of hepatic transaminase—In placebo-controlled schizophrenia trials, clinically significant ALT (SGPT) elevations (≥3 times the upper limit of the normal range) were observed in 2% (6/243) of patients exposed to ZYPREXA compared to none (0/115) of the placebo patients. None of these patients developed jaundice. 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. Periodic assessment of transaminases is recommended in patients with significant hepatic disease.

Special populations, elderly—Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Olanzapine should be used with caution in patients at risk for aspiration pneumonia. In 5 studies in elderly patients with dementia-related psychosis, adverse events reported more commonly with olanzapine than with placebo were falls, somnolence, peripheral edema, abnormal gait, urinary incontinence, lethargy, increased weight, asthenia, pyrexia, pneumonia, dry mouth, and visual hallucinations. Olanzapine should be used with caution in elderly patients with dementia. Olanzapine is not approved for treatment of patients with dementia-related psychosis.

Drug interactions—Coadministration of diazepam or ethanol with ZYPREXA may potentiate orthostatic hypotension. Lower doses of ZYPREXA should be considered in patients receiving concomitant therapy with fluvoxamine.

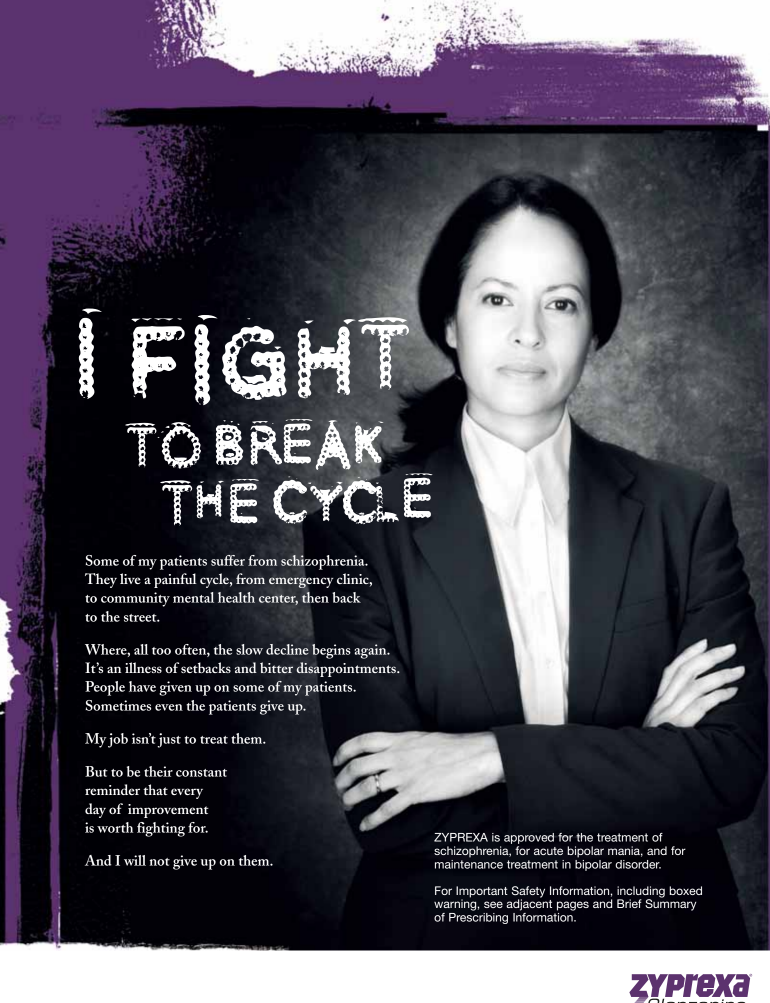
Medication dispensing and prescribing errors have occurred between ZYPREXA® (olanzapine) and Zyrtec® (cetirizine HCl). These errors could result in unnecessary adverse events or potential relapse in patients suffering from schizophrenia or bipolar disorder. To reduce the potential for dispensing errors,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

The most common treatment-emergent adverse events associated with ZYPREXA (vs placebo) in 6-week acute-phase schizophrenia trials were somnolence (26% vs 15%), 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 events associated with ZYPREXA (vs placebo) in 3- and 4-week bipolar mania trials were somnolence (35% vs 13%), 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%).

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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: 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 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: 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. ZYPREXA is not approved for the treatment of patients with dementia-related psychosis (see BOX WARNING).

psychosis (see BOX WARNING).

In placebo-controlled clinical trials of elderly patients with dementia-related psychosis, the incidence of death in olanzapine-treated patients (3.5%) was significantly greater than placebo-treated patients (1.5%).

Cerebrovascular Adverse Events. Including Stroke, in Elderly Patients with Dementia—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 higher incidence of cerebrovascular adverse events in patients treated with olanzapine compared to patients treated with placebo. Olanzapine is not approved for the treatment of patients with dementia-related psychosis.

Hyperplusepia and Displace Molitive—Hyperplusepia in some access associated with kalazaidacis come of

<u>Hyperglycemia and Diabetes Mellitus</u>—Hyperglycemia, 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 increased background risk of diabetes in patients with schizophrenia and the increasing incidence of diabetes mellitus in the general population. Patients diagnosed with diabetes who are started on atypical antipsychotics should be monitored 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

starting treatment. Any patient treated with atypicals should be monitored for symptoms of hyperglycemia. Patients who develop symptoms of hyperglycemia during treatment with atypicals should undergo FBG testing. Neuroleptic Malignant Syndrome (NMS)—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>Tardive Dyskinesia (TD)</u>—Potentially irreversible TD may develop in patients treated with antipsychotic drugs. Although the prevalence of TD appears to be highest among the elderly, especially elderly women, it is impossible to predict which patients are more likely to develop the syndrome. If signs and symptoms of TD appear, consider drug discontinuation.

PRECAUTIONS: Hemodynamic Effects—Olanzapine 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 olanzapine for injection. Incidence of syncope was 0.6%, 15/2500 with oral olanzapine in phase 2-3 trials and 0.3%, 2/722 with intramuscular olanzapine for injection in clinical trials. Three normal volunteers in phase 1 studies with intramuscular olanzapine experienced hypotension, bradycardia, and sinus pauses of up to 6 seconds that spontaneously resolved (in 2 cases the events occurred on intramuscular olanzapine, and in 1 case, on oral sportlaneously resolved (III 2 cases the events occurred on intrahiuscular oralization, and in 1 case, on oldarapine). 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 drows or dizzy after injection with intramuscular olanzapine for injection until examination has indicated they are not experiencing postural hypotension, bradycardia, and/or hypoventilation. Olanzapine has indicated they are not experiencing postural hypotension, bradycardia, and/or hypoventilation. Juarapine should be used with particular caution in patients with known cardiovascular disease, (history of myocardial 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 put them at increased medical risk. Caution is necessary in patients receiving treatment with other drugs having effects that can induce hypotension, bradycardia, respiratory or CNS depression (see Drug Interactions). Concomitant administration of intramuscular olanzapine and parenteral benzodiazepine has not been studied and is not recommended. If such combination treatment is considered, careful evaluation of clinical status for expessive extation and cardinesen/status (hepression) is recommended.

and is not recommended, it such combination treather its considered, careful evaluation of clinical status in excessive sedation and cardiorespiratory depression is recommended.

<u>Seizures</u>—During premarketing testing, 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.

Hyperprolactinemia—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.

tumorigenesis in humans; the available evidence is inconclusive.

<u>Transaminase Elevations</u>—In placebo-controlled studies, clinically significant ALT (SGPT) elevations (≥3 times the upper limit of normal) were observed in 2% (6/243) of patients exposed to olanzapine compared to no (0/115) placebo patients. None of these patients experienced jaundice. Among about 2400 patients with baseline SGPT ≥90 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 olanzapine 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 paried. Exercise eaution in natients who have since and symptomes of heaptic impairment. 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).

Potential for Cognitive and Motor Impairment—Somnolence was a commonly reported, dose-related adverse event in premarketing trials (olanzapine 26% vs placebo 15%). Somnolence led to discontinuation in 0.4%

(9/2500) of patients in the oral premarketing database.

Body_Temperature_Regulation—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 mortality in patients with advanced Alzheimer'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

Suicide—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 olarazapine should be written for the smallest quantity of tablets consistent with good patient management.

<u>Use in Patients with Concomitant Illnesses</u>—Olanzapine should be used with caution in patients with clinically significant prostatic hypertrophy, narrow angle glaucoma, or a history of paralytic 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 than with placebo: falls, somnolence, peripheral edema, abnormal gait, urinary incontinence, lethargy, increased weight, asthenia, pyrexia, pneumonia, dry mouth, visual hallucinations. Discontinuation due adverse events was significantly greater with olanzapine than placebo (13% vs 7%). Elderly patients with dementia-related psychosis treated with olanzapine are at an increased risk of death compared to placebo.

Olanzapine is not approved for treatment of patients with dementia-related psychosis. If the prescriber elects to treat this patient population, vigilance should be exercised (see BOX WARNING and WARNINGS).

Because of the risk of orthostatic hypotension with olanzapine, use caution in cardiac patients (see

Hemodynamic Effects)

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

<u>Laboratory Tests</u>—Periodic assessment of transaminases is recommended in patients with significant

<u>Drug Interactions</u>—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 enzyme (e.g., 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 inhibition 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 C_{\max} 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. Carbamazepine (200 mg bid) causes an approximately 50% increase in the clearance of of olarzapine. Higher daily doses of carbamazepine may cause an experimental of the clearance of olarzapine. Higher daily doses of carbamazepine may cause an even greater increase in olarzapine clearance. Neither ethanol (45 mg/70 kg single dose) nor warfarin (20 mg single dose) had an effect on olarzapine pharmacokinetics. Fluoxetine at 60 mg (single or multiple doses) causes a small increase in the C_{max} of olarzapine and a small decrease in olarzapine clearance; however, the impact of this factor is small in comparison to the overall variability between individuals, and dose modification is not routinely recommended. Fluoxamine decreases the clearance of olarzapine; lower doses of olarzapine should be

recommended. Fluvoxamine decreases the clearance of olanzapine; lower doses of olanzapine should be considered in patients receiving fluvoxamine concomitantly. In vitro data suggest that a clinically significant pharmacokinetic interaction between olanzapine and valproate is unlikely.

Olanzapine is unlikely to cause clinically important drug interactions mediated by the enzymes CYP1A2, CYP2C9, CYP2C19, CYP2C9, CYP2C9A. Single doses of olanzapine did not affect the pharmacokinetics of imipramine/desipramine or warfarin. Multiple doses of olanzapine did not influence the kinetics of diazepam/N-desmethyldiazepam, lithium, ethanol, or biperiden. However, coadministration of either diazepam or ethanol potentiated the orthostic hypotension observed with olanzapine. Multiple doses of olanzapine did not affect the pharmacokinetics of theophylling or its metabolities. Coadministration of internavely large agent. not affect the pharmacokinetics of theophylline or its metabolites. Co-administration of intramuscular lorazepam and intramuscular olanzapine for injection added to the somnolence observed with either drug alone (see Hemodynamic Effects).

Hemodynamic Effects).

<u>Carcinogenesis. Mutagenesis. Impairment of Fertility</u>—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 males in the 30/20 mg/kg/d group. The incidence of mammary gland adenomas and adenocarcinomas was significantly increased in female mice and rats 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-fold 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). Diestrous was prolonged and estrous delayed at 0.6 times the MHDOD (mg/m² basis); therefore, olanzapine may produce a delay in ovulation.

Pregnancy Category C—There are no adequate and well-controlled studies in pregnant women. Olanzapine

basis); Intercitore, danazapine may produce a delay in ovolation.

<u>Pregnancy Category C</u>—There are no adequate and well-controlled studies in pregnant women. Olanzapine should be used in pregnancy only if the potential benefit justifies the potential risk to the fetus.

<u>Labor and Delivery, Nursing Mothers</u>—Parturition in rats was not affected by olanzapine; its effect on labor and delivery in humans is unknown. In a study in lactating, healthy women, olanzapine was excreted in breast milk. Mean infant dose at steady state was estimated to be 1.8% of the maternal dose. It is recommended that

women receiving olanzapine should not breast-feed.

<u>Use in Pediatric and Geriatric Patients</u>—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 established. In premarketing clinical trails in patients with schizophrenia, there was no indication of any different tolerability of olanzapine in the elderly compared to younger patients. Studies in elderly patients with dementia-related psychosis have suggested there may be a different tolerability profile in these patients. Elderly patients with dementia-related psychosis treated with olanzapine are at an increased risk of death compared to placebo. Olanzapine is not approved for treatment of patients with dementia-related psychosis. If the prescriber elects to treat these patients, vigilance should be exercised. Consider a lower starting dose for any geriatric patient in the presence of factors that might decrease pharmacokinetic clearance or increase the pharmacodynamic response to olanzapine (see BOX WARNING, WARNINGS, and PRECAUTIONS).

to olarzapine (see BOX WARNING, WARNINGS, and PRECAUTIONS).

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 olarzapine and 722 patients with exposure to intramuscular olarzapine for injection, including patients with schizophrenia, bipolar mania, or Alzheimer's disease (oral olarzapine trials) and patients with agitation associated with schizophrenia, bipolar I disorder (manic or mixed episodes), or dementia (intramuscular olarzapine 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 gain, laboratory changes, and ECG 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 olarzapine rusia (olanzapine pus lithium or valproate) so 2%; bipolar mania cotherapy, 11% [olanzapine plus lithium or valproate alone]); or in placebo-controlled intramuscular olanzapine for injection trials (olanzapine for injection, 0.4%; 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

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 ≥5% and olanzapine incidence at least twice that for placebo) were: postural hypotension, constipation, weight gain, dizziness, personality disorder (COSTART term for nonaggressive 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, somnolence, dizziness, and tremor. In short-term bipolar mania combination therapy trials, the most common treatment-emergent adverse events observed with olaryonia pulse lithium or valorate were due. common treatment-emergent adverse events observed with olanzapine plus lithium or valproade were dry mouth, weight gain, increased appetite, dizziness, back pain, constipation, speech disorder, increased salivation, amnesia, and paresthesia. In 24-hour placebo-controlled trials of intramuscular olanzapine for injection for

alments, and parestress. III ≥4-nour placeu0-continued untal of intrainscular oralization is observed at an incidence of ≤5% and at least twice that for placebo (olanzapine for injection 6%, placebo 3%). Adverse Events with an Incidence ≥2% in Oral Monotherapy Trials—The following treatment-emergent events were reported at an incidence of ≥2% with oral olanzapine (doses ≥2.5 mg/d), and at a greater incidence with olanzapine than with placebo in short-term placebo-controlled trials (olanzapine N=532, placebo N=294): Body as a Whole—accidental injury, asthenia, fever, back pain, chest pain; Cardiovascular—postural hypotension, tachycardia, hypertension; Digestive—dry mouth, constipation, dyspepsia, vomiting, increased appetite; Hemic and Lymphatic—ecchymosis; Metabolic and Nutritional—weight gain, peripheral edema;

appettie. **Hemic and Lymphatic**—ecchymosis; **Metabolic and Nutritional**—weight gain, peripheral edema; **Musculoskeletal**—extremity pain (other than joint), joint pain; **Nervous System**—somnolence, insominidizines, abnormal gait, tempor, akthisia, hypertonia, articulation impairment; **Respiratory**—ininitis, cough increased, pharyngitis; **Special Senses**—amblyopia; **Urogenital**—urinary incontinence, urinary tract infection. **Adverse Events with an Incidence \$2.2% in Oral Combination Therapy Trials**—The following treatment emergent events were reported at an incidence of \$2.7% with oral olarazpine (doses \$5 mg/d) plus lithium or valproate (N=229), and at a greater incidence than with placebo plus lithium or valproate (N=115) in short-term placebo-controlled trials: **Body as a Whole**—asthenia, back pain, accidental injury, chest pain; **Cardiovascular**—hypertension; **Digestive**—dry mouth, increased appetite, thirst, constipation, increased salivation; **Metabolic and Nutritional**—weight gain, peripheral edema, edema; **Nervous System**—somnolence, termor, depression, dizziness, speech disorder, amnesia, paresthesia, apathy, confusion, euphoria, incoordination; **Respiratory**—pharyngitis, dyspnea; **Skin and Appendages**—sweating, acne, dry skin; **Special Senses**—amblyopia, abnormal vision; **Urogenital**—dysmenorrhea, vaginitis.**
**Adverse Events with an Incidence of \$1%* with intramuscular rolas="The following treatment-emergent adverse events were reported at an incidence of \$1%* with intramuscular rolas="The following treatment-emergent adverse events were reported at an incidence of \$1.5%* with intramuscular rolas="The following treatment-emergent adverse events were reported at an incidence of \$1.5%* with intramuscular rolas="The following treatment-emergent adverse events were reported at an incidence of \$1.5%* with intramuscular rolas="The following treatment-emergent adverse events were reported at an incidence of \$1.5%* with intramuscular olanzapine for injection (2.5-10 mg/inject

schizophrenia or bipolar mania: **Body as a Whole**—asthenia; **Cardiovascular**—hypotension, postural hypotension; **Nervous System**—somnolence, dizziness, tremor.

PV 5195 AMP

<u>Dose Dependency of Adverse Events in Short-Term. Placebo-Controlled Trials—Extrapyramidal Symptoms—</u>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 akathisia events (spontaneously reported COSTART Akadinsk global score 22.1 int leastife fliat, villy akadinsk events (sportinedusty) reported COSTAM terms akathisk 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 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 spontaneously reported adverse events.

Other Adverse Events—Dose-relatedness of adverse events was assessed using data from a clinical

trial involving 3 fixed oral dosage ranges compared with placebo. The following treatment-emergent events showed a statistically significant trend: asthenia, dry mouth, nausea, somnolence, tremor.

<u>Vital Sign Changes</u>—Oral olanzapine was associated with orthostatic hypotension and tachycardia in clinical trials. Intramuscular olanzapine for injection was associated with bradycardia, hypotension, and tachycardia in clinical trials (see PRECAUTIONS).

tachycardia in clinical trials (see PRECAUTIONS).

Weight Gain—In placebo-controlled 6-week schizophrenia studies, weight gain was reported in 5.6% of oral olanzapine patients (average 2.8-kg gain) compared to 0.8% of placebo patients (average 0.4-kg loss); 29% of olanzapine patients gained >7% of their baseline weight, compared to 3% of placebo patients. During continuation therapy (238 median days of exposure), 56% of patients met the criterion for having gained >7% of their baseline weight. Average gain during long-term therapy was 54 kg. Laboratory Changes—Olanzapine is associated with asymptomatic increases in SGPT, SGOT, and GGT and with increases in serum prolactin and CPK (see PRECAUTIONS). Asymptomatic elevation of eosinophils was reported in 0.3% of olanzapine patients in premarketing trials. There was no indication of a rick of clinically significant neutronegia associated with plazazapine in the programacyting database.

of a risk of clinically significant neutropenia associated with olanzapine in the premarketing database.

In clinical trials among olanzapine-treated patients with baseline random triglyceride levels of <150 mg/dL (N=659), 0.5% experienced triglyceride levels of ≥500 mg/dL anytime during the trials. In <150 mg/dL (N=659), 0.5% experienced trighyceride levels of ≥500 mg/dL anytime during the trals. In these same trials, olanzapine-treated patients (N=1185) had a mean trighyceride increase of 20 mg/dL from a mean baseline of 175 mg/dL. In placebo-controlled trials, olanzapine-treated patients with baseline random cholesterol levels of <200 mg/dL (N=1034) experienced cholesterol levels of ≥240 mg/dL anytime during the trials more often than placebo-treated patients (N=062; 3.6% vs. 2.2% respectively). In these same trials, olanzapine-treated patients (N=2528) had a mean increase of 0.4 mg/dL in cholesterol from 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.</p>

baseline of 203 mg/dl.

<u>ECG Changes</u>—Analyses of pooled placebo-controlled trials revealed no statistically significant olanzapine/placebo differences in incidence of potentially important changes in ECG parameters, including QT, QTc, and PR intervals. Olanzapine was associated with a mean increase in heart rate of 2.4 BPM compared to no change among placebo patients.

<u>Other Adverse Events Observed During Clinical Trials</u>—The following treatment-emergent events were reported with oral olanzapine at multiple doses ≥1 mg/d in clinical trials (8661 patients, 4165 patient-years of exposure). This list may not include events previously listed elsewhere in labeling those events for which a drug cause was remote. These terms which were so general as be be

4165 patient-years of exposure). 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. Frequent events occurred in ≥1/100 patients; infrequent events occurred in 1/100 to 1/1000 patients; rare events occurred in ≥1/1000 patients. Bady as a Whole—Frequent: dental pain, flu syndrome; Infrequent: abdomen enlarged, chills, face edema, intentional injury, malaise, moniliasis, neck pain, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt; Rare: chills and fever, hangover effect, sudden death. Cardiovascular—Frequent: hypotension; Infrequent: atrial fibrillation, bradycardia, cerebrovascular accident, congestive heart failure, heart arrest hemorrhage migraine pallor, abilistion, vascodilation, venticular extrasystoles. Pages injury, malaise, moniliasis, neck pain, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt; **Rare: chills and fever, hangover effect, sudden death. **Cardiovascular—Frequent: hypotension, Infrequent: atrial fibrillation, bradycardia, cerebrovascular accident, congestive heart failure, pelmonary embolis. **Digestive—Frequent: talkulene, increased salivation, thirst: **Infrequent: dysphagia, esophagitis, fecal impaction, fecal incontinence, gastritis, gastroenteritis, gingivitis, hepatitis, melena, mouth ulceration, nausea and vomiting, oral moniliasis, periodontal abscess, rectal hemorrhage, stomatitis, tongue edema, tooth caries; **Rare: aphthous stomatitis, entertiis, eructation, esophageal ulcer, glossitis, fleus, intestinal obstruction, fiver fatty deposit, tongue discoloration. **Endocrine—Infrequent: diabetes mellitus; **Rare: diabetic acidosis, goiter. **Hemic and Lymphatic—Infrequent: anemia, cyanosis, leukocytosis, leukopenia, lymphadenopathy, thrombocytopenia, **Rare: normocytic anemia, thrombocythemia, Metabolic and Nutritional—Infrequent: acidosis, alkaline phosphatase increased, bilirubinemia, dehydration, hypercholesteremia, hyperplyemia, hyperuricemia, hypoglyeemia, hypokalemia, hyponatremia, lower extremity edema, upper extremity edema; **Rare: gout, hyperkalemia, hyporatemia, hyporosis, rhematodia arthritis. **Nervous System—Frequent: abnormal dreams, amnesia, delusions, emotional lability, euphoria, manic reaction, paresthesia, schizophrenic reaction, infrequent: akinga, alcohol misuse, antisocial reaction, ataxia, CNS stimulation, cogwheel rigidity, delirium, dementia, depersonalization, dysarthria, facial paralysis, hyposthesia, hypokinesia, hypotonia, iacorodination, libido increased, obsessive compulsive symptoms, phobias, somatization, stimulant misuse, stupor, stuttering, tardive dyskinesia, veritgo, withdrawal syndrome; **Rare: acidonation, misuse, antisocial reaction, glaucoma, kerzoba, propositis lens. **Urgenital—Frequent: wagnitis**, Infrequent: abnormal pacia, t

DRUG ABUSE AND DEPENDENCE: Olanzapine is not a controlled substance.

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Literature revised March 20, 2006



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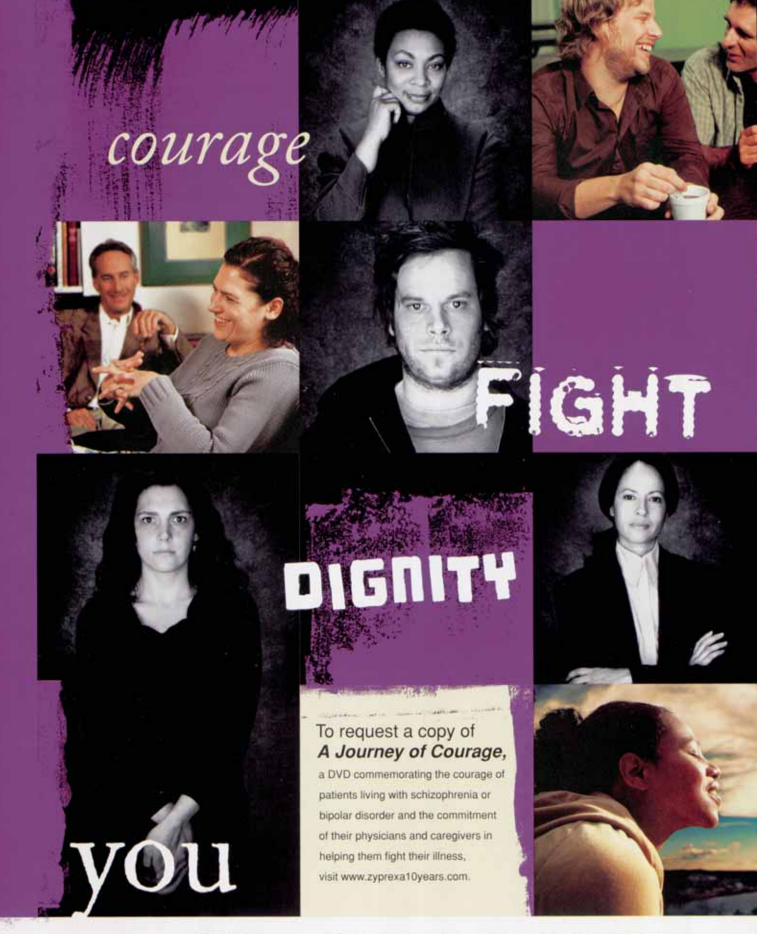
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10332E-10-06

Success isn't just measured in years, it's measured in moments.





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

See under flap on adjacent page for Important Safety Information, including boxed warning, and Brief Summary of Prescribing Information.



We celebrate the moments of the last 10 years

that you've helped make possible.

Along with the 150,000 physicians¹ who last year prescribed ZYPREXA, Lilly is proud to be a part of the ongoing fight to make a difference in patients' lives.

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.



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 drugtreated 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 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 elderly 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.

Tardive dyskinesia (TD)—As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of TD. If its signs and symptoms appear, discontinuation should be considered.

Orthostatic hypotension—In premarketing schizophrenia trials, some patients taking ZYPREXA may have experienced orthostatic hypotension associated with dizziness, tachycardia, and, in some cases, syncope (15/2500, 0.6%).

Seizures—Occurred infrequently in premarketing clinical trials (22/2500, 0.9%). ZYPREXA should be used cautiously in patients with a history of seizures or with conditions that lower the seizure threshold.

Effect on prolactin—Modest elevations of prolactin were seen with ZYPREXA in acute-phase schizophrenia trials (incidence 34% vs 13% with placebo), although mean changes from baseline to endpoint were not statistically significantly different between olanzapine and placebo. Some patients may have persisting modest prolactin elevations.

Transient, asymptomatic elevations of hepatic transaminase—
In placebo-controlled schizophrenia trials, clinically significant ALT
(SGPT) elevations (≥3 times the upper limit of the normal range)
were observed in 2% (6/243) of patients exposed to ZYPREXA
compared to none (0/115) of the placebo patients. None of these
patients developed jaundice. 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. Periodic
assessment of transaminases is recommended in patients with
significant hepatic disease.

Special populations, elderly—Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Olanzapine should be used with caution in patients at risk for aspiration pneumonia. In 5 studies in elderly patients with dementia-related psychosis, adverse events reported more commonly with olanzapine than with placebo were falls, somnolence, peripheral edema, abnormal gait, urinary incontinence, lethargy, increased weight, asthenia, pyrexia, pneumonia, dry mouth, and visual hallucinations. Olanzapine should be used with caution in elderly patients with dementia. Olanzapine is not approved for treatment of patients with dementia-related psychosis.

Drug interactions—Coadministration of diazepam or ethanol with ZYPREXA may potentiate orthostatic hypotension. Lower doses of ZYPREXA should be considered in patients receiving concomitant therapy with fluvoxamine.

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 events associated with ZYPREXA (vs placebo) in 6-week acute-phase schizophrenia trials were somnolence (26% vs 15%), 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 events associated with ZYPREXA (vs placebo) in 3- and 4-week bipolar mania trials were somnolence (35% vs 13%), 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%).

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IMS Health, National Prescription Audit Plus™, Jan-Dec 2005, ZYP20060810A.

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<u>Labor and Delivery. Nursing Mothers</u>—Parturition in rats was not affected by olanzapine; its effect on labor and delivery in humans is unknown. In a study in lactating, healthy women, olanzapine was excreted in breast milk. Mean infant dose at steady state was estimated to be 1.8% of the maternal dose. It is recommended that women receiving olanzapine should not breast-feed.

Use in Pediatric and Geriatric 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 olanzapine in the elderly compared to younger patients. Studies in elderly patients with dementia-related psychosis have suggested there may be a different tolerability profile in these patients. Elderly patients with dementia-related psychosis treated with olanzapine are at an increased risk of death compared to placebo. Olanzapine is not approved for treatment of patients with dementia-related psychosis. If the prescriber elects to treat these patients, vigilance should be exercised. Consider a lower starting dose for any geriatric patient in the presence of factors that might decrease pharmacokinetic clearance or increase the pharmacodynamic response to olanzapine (see BOX WARNING, WARNINGS, and PRECAUTIONS).

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 (oral olanzapine trials) and patients with agitation associated with schizophrenia, bipolar I disorder (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 gain, laboratory changes, and ECG 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 monotherapy, 2% vs 2%; bipolar mania cotherapy, 11% [olanzapine plus lithium or valproate] vs 2% [lithium or valproate alone]); or in placebo-controlled intramuscular olanzapine for injection trials (olanzapine for injection, 0.4%; 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 ≥5% and olanzapine incidence at least twice that for placebo) were: postural hypotension, constipation, weight gain, dizziness, personality disorder (COSTART term for nonaggressive 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, somnolence, dizziness, and tremor. In short-term bipolar mania combination therapy trials, the most common treatment-emergent adverse events observed with olanzapine plus lithium or valproate were dry mouth, weight gain, increased appetite, dizziness, back pain, constipation, speech disorder, increased salivation, amnesia, and paresthesia. In 24-hour placebo-controlled trials of intramuscular olanzapine for injection for agitation associated with schizophrenia or bipolar mania, somnolence was the one adverse event observed at an incidence of ≥5% and at least twice that for placebo (olanzapine for injection 6%, placebo 3%).

Adverse Events with an Incidence ≥2% in Oral Monotherapy Trials—The following treatmentemergent events were reported at an incidence of ≥2% with oral olanzapine (doses ≥2.5 mg/d), and at
a greater incidence with olanzapine than with placebo in short-term placebo-controlled trials
(olanzapine N=532, placebo N=294): Body as a Whole—accidental injury, asthenia, fever, back pain,
chest pain; Cardiovascular—postural hypotension, tachycardia, hypertension; Digestive—dry
mouth, constipation, dyspepsia, vomiting, increased appetite; Hemic and Lymphatic—ecchymosis;
Metabolic and Nutritional—weight gain, peripheral edema; Musculoskeletal—extremity pain (other
than joint), joint pain; Nervous System—somnolence, insomnia, dizziness, abnormal gait, tremor,
akathisia, hypertonia, articulation impairment; Respiratory—rhinitis, cough increased, pharyngitis;
Special Senses—amblyopia; Urogenital—urinary Incontinence, urinary tract infection.

Special Senses—amblyopia; Urogenital—urinary incontinence, urinary tract infection.

Adverse Events with an Incidence ≥% in Oral Combination Therapy Trials—The following treatment-emergent events were reported at an incidence of ≥2% with oral olanzapine (doses ≥5 mg/d) plus lithium or valproate (N=229), and at a greater incidence than with placebo plus lithium or valproate (N=115) in short-term placebo-controlled trials: Body as a Whole—asthenia, back pain, accidental injury, chest pain; Cardiovascular—hypertension; Digestive—dry mouth, increased appetite, thirst, constipation, increased salivation; Metabolic and Nutritional—weight gain, peripheral edema, edema; Nervous System—somnolence, tremor, depression, dizziness, speech disorder, amnesia, paresthesia, apathy, confusion, euphoria, incoordination; Respiratory—pharyngitis, dyspnea; Skin and Appendages—sweating, acne, dry skin; Special Senses—amblyopia, abnormal vision; Urogenital—dysmenorrhea, vaginitis.

Adverse Events with an Incidence ≥1% in Intramuscular Trials—The following treatment-emergent adverse events were reported at an incidence of ≥1% with intramuscular olanzapine for injection (2.5 –10 mg/injection) 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, 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 akathisia events (spontaneously reported 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 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 spontaneously reported adverse events.

Other Adverse Events—Dose-relatedness of adverse events was assessed using data from a clinical trial involving 3 fixed oral dosage ranges compared with placebo. The following treatment-emergent events showed a statistically significant trend: asthenia, dry mouth, nausea, somnolence, tremor.

<u>Vital Sign Changes</u>—Oral olanzapine was associated with orthostatic hypotension and tachycardia in clinical trials. Intramuscular olanzapine for injection was associated with bradycardia, hypotension, and tachycardia in clinical trials (*see* PRECAUTIONS).

Weight Gain—In placebo-controlled 6-week schizophrenia studies, weight gain was reported in 5.6% of oral olanzapine patients (average 2.8-kg gain) compared to 0.8% of placebo patients (average 0.4-kg loss); 29% of olanzapine patients gained >7% of their baseline weight, compared to 3% of placebo patients. During continuation therapy (238 median days of exposure), 56% of patients met the criterion for having gained >7% of their baseline weight. Average gain during long-term therapy was 5.4 kg.

<u>Laboratory Changes</u>—Olanzapine is associated with asymptomatic increases in SGPT, SGOT, and GGT and with increases in serum prolactin and CPK (*see* PRECAUTIONS). Asymptomatic elevation of eosinophils was reported in 0.3% of olanzapine patients in premarketing trials. There was no indication of a risk of clinically significant neutropenia associated with olanzapine in the premarketing database.

In clinical trials among olanzapine-treated patients with baseline random triglyceride levels of <150 mg/dL (M=659), 0.5% experienced triglyceride levels of \geq 500 mg/dL anytime during the trials. In these same trials, olanzapine-treated patients (M=1185) had a mean triglyceride increase of 20 mg/dL from a mean baseline of 175 mg/dL. In placebo-controlled trials, olanzapine-treated patients with baseline random cholesterol levels of <200 mg/dL (M=1034) experienced cholesterol levels of M=240 mg/dL anytime during the trials more often than placebo-treated patients (M=602; 3.6% vs 2.2% respectively). In these same trials, olanzapine-treated patients (M=628) had a mean increase of 0.4 mg/dL in cholesterol from a mean baseline of 203 mg/dL, which was significantly different compared to placebo-treated patients (M=1415) with a mean decrease of 4.6 mg/dL from a mean baseline of 203 mg/dL.

<u>ECG Changes</u>—Analyses of pooled placebo-controlled trials revealed no statistically significant olanzapine/placebo differences in incidence of potentially important changes in ECG parameters, including QT, QTc, and PR intervals. Olanzapine was associated with a mean increase in heart rate of 2.4 BPM compared to no change among placebo patients.

Other Adverse Events Observed During Clinical Trials—The following treatment-emergent events were reported with oral olanzapine at multiple doses ≥1 mg/d in clinical trials (8661 patients, 4165 patient-years of exposure). 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. Frequent events occurred in ≥1/100 patients; infrequent events occurred in 1/100 to 1/1000 patients; rare events occurred in <1/1000 patients. Body as a Whole-Frequent: dental pain, flu syndrome; Infrequent: abdomen enlarged, chills, face edema, intentional injury, malaise, moniliasis, neck pain, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt; Rare: chills and fever, hangover effect, sudden death. Cardiovascular—Frequent: hypotension; Infrequent: atrial fibrillation, bradycardia, cerebrovascular accident, congestive heart failure, heart arrest, hemorrhage, migraine, pallor, palpitation, vasodilatation, ventricular extrasystoles; Rare: arteritis, heart failure, pulmonary embolus. Digestive—Frequent: flatulence, increased salivation, thirst; Infrequent: dysphagia, esophagitis, fecal impaction, fecal incontinence, gastritis, gastroenteritis, gingivitis, hepatitis, melena, mouth ulceration, nausea and vomiting, oral moniliasis, periodontal abscess, rectal hemorrhage, stomatitis, tongue edema, tooth caries; Rare: aphthous stomatitis, enteritis, eructation, esophageal ulcer, glossitis, ileus, intestinal obstruction, liver fatty deposit, tongue discoloration. Endocrine Infrequent: diabetes mellitus; Rare: diabetic acidosis, goiter. Hemic and Lymphatic—Infrequent: anemia, cyanosis, leukocytosis, leukopenia, lymphadenopathy, thrombocytopenia; Rare: normocytic anemia, thrombocythemia. Metabolic and Nutritional-Infrequent: acidosis, alkaline phosphatase increased, bilirubinemia, dehydration, hypercholesteremia, hyperglycemia, hyperlipemia, hyperuricemia, hypoglycemia, hypokalemia, hyponatremia, lower extremity edema, upper extremity edema; *Rare*: gout, hyperkalemia, hypernatremia, hypoproteinemia, ketosis, water intoxication. *Musculoskeletal—Frequent:* joint stiffness, twitching; *Infrequent:* arthritis, arthrosis, leg cramps, myasthenia; Rare: bone pain, bursitis, myopathy, osteoporosis, rheumatoid arthritis. Nervous System-Frequent: abnormal dreams, amnesia, delusions, emotional lability, euphoria, manic reaction, paresthesia, schizophrenic reaction; Infrequent: akinesia, alcohol misuse, antisocial reaction, ataxia, CNS stimulation, cogwheel rigidity, delirium, dementia, depersonalization, dysarthria, facial paralysis, hypesthesia, hypokinesia, hypotonia, incoordination, libido decreased, libido increased, obsessive compulsive symptoms, phobias, somatization, stimulant misuse, stupor, stuttering, tardive dyskinesia, vertigo, withdrawal syndrome; Rare: circumoral paresthesia, coma, encephalopathy, neuralgia, neuropathy, nystagmus, paralysis, subarachnoid hemorrhage, tobacco misuse. Respiratory-Frequent: dyspnea; Infrequent: apnea, asthma, epistaxis, hemoptysis, hyperventilation, hypoxia, laryngitis, voice alteration; Rare: atelectasis, hiccup, hypoventilation, lung edema, stridor. Skin and Appendages-Frequent: sweating; Infrequent: alopecia, contact dermatitis, dry skin, eczema, maculopapular rash, pruritus, seborrhea, skin discoloration, skin ulcer, urticaria, vesiculobullous rash; Rare: hirsutism, pustular rash. Special Senses—Frequent: conjunctivitis; Infrequent: abnormality of accommodation, blepharitis, cataract, deafness, diplopia, dry eyes, ear pain, eye hemorrhage, eye inflammation, eye pain, ocular muscle abnormality, taste perversion, tinnitus; Rare: corneal lesion, glaucoma, keratoconjunctivitis, macular hypopigmentation, miosis, mydriasis, pigment deposits lens. *Urogenital—Frequent*, relation, "mosts, machinal repopulation," amenorrhea, "breast pain, cystitis, decreased menstruation," dysuria, female lactation," glycosuria, gynecomastia, hematuria, impotence, * increased menstruation, * menorrhagia, * metrorrhagia, * polyuria, premenstrual syndrome,* pyuria, urinary frequency, urinary retention, urinary urgency, urination impaired, uterine fibroids enlarged, * vaginal hemorrhage*; Rare: albuminuria, breast enlargement, mastitis, oliguria. (*Adjusted for gender.)

The following treatment-emergent events were reported with intramuscular olanzapine for injection at one or more doses ≥2.5 mg/injection 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 events 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; Infrequent: abdominal pain, fever. Cardiovascular—Infrequent: AV block, heart block, syncope. Digestive—Infrequent: diarrhea, nausea. Hemic and Lymphatic—Infrequent: anemia. Metabolic and Nutritional—Infrequent: creatine phosphokinase increased, dehydration, hyperkalemia. Musculoskeletal—Infrequent: twitching. Nervous System—Infrequent: abnormal gait, akathisia, articulation impairment, confusion, emotional lability. Skin and Appendages—Infrequent: sweating.

Postintroduction Reports—Reported since market introduction and temporally (not necessarily causally) related to olanzapine therapy: allergic reaction (eg, anaphylactoid reaction, angioedema, pruritus or urticaria), diabetic coma, jaundice, pancreatitis, priapism, rhabdomyolysis, and venous thromboembolic events (including pulmonary embolism and deep venous thrombosis). Random cholesterol levels of ≥240 mg/dL and random triglyceride levels of ≥1000 mg/dL have been rarely reported.

DRUG ABUSE AND DEPENDENCE: Olanzapine is not a controlled substance.

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Literature revised March 20, 2006

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ZYPREXA® ZYDIS® Olanzapine Orally Disintegrating Tablets

ZYPREXA® IntraMuscular Olanzapine for Injection

Brief Summary: Please consult package insert for complete prescribing information.

WARNING: 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 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: 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. 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 olanzapine-treated patients (3.5%) was significantly greater than placebo-treated patients (1.5%).

Cerebrovascular Adverse Events. Including Stroke, in Elderly Patients with Dementia—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 higher incidence of cerebrovascular adverse events in patients treated with olanzapine compared to patients treated with placebo. Olanzapine 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 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 with schizophrenia and the increasing incidence of diabetes mellitus in the general population. Patients diagnosed with diabetes who are started on atypical antipsychotics should be monitored 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 atypicals should be monitored for symptoms of hyperglycemia. Patients who develop symptoms of hyperglycemia during treatment with atypicals should undergo FBG testing.

Neuroleptic Malignant Syndrome (NMS)—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.

Tardive Dyskinesia (TD)—Potentially irreversible TD may develop in patients treated with antipsychotic drugs. Although the prevalence of TD appears to be highest among the elderly, especially elderly women, it is impossible to predict which patients are more likely to develop the syndrome. If signs and symptoms of TD appear, consider drug discontinuation.

PRECAUTIONS: <u>Hemodynamic Effects</u>—Olanzapine 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 clanzapine for injection. Incidence of syncope was 0.6%, 15/2500 with oral olanzapine in phase 2-3 trials and 0.3%, 2/722 with intramuscular olanzapine for injection in clinical trials. Three normal volunteers in phase 1 studies with intramuscular olanzapine experienced hypotension, bradycardia, and sinus pauses of up to 6 seconds that spontaneously resolved (in 2 cases the events occurred on intramuscular olanzapine, and in 1 case, on oral olanzapine). 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 dizzy after injection with intramuscular olanzapine 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 myocardial 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 put them at increased medical risk. Caution is necessary in patients receiving treatment with other drugs having effects that can induce hypotension, bradycardia, respiratory or CNS depression (see Drug Interactions). Concomitant administration of intramuscular olanzapine and parenteral benzodiazepine has not been studied and is not recommended. If such combination treatment is considered, careful evaluation of clinical status for excessive sedation and cardiorespiratory depression is recommended.

<u>Seizures</u>—During premarketing testing, seizures occurred in 0.9% (22/2500) of olanzapinetreated patients, 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.

Transaminase Elevations—In placebo-controlled studies, clinically significant ALT (SGPT) elevations (\$\approx 3\$ times the upper limit of normal) were observed in 2% (6/243) of patients exposed to olanzapine compared to no (0/115) placebo patients. None of these patients experienced jaundice. Among about 2400 patients with baseline SGPT \$\approx 90\$ IU/L, 2% (50/2381) had asymptomatic SGPT elevations to \$\approx 20\$ IU/L. Most were transient changes that tended to normalize while olanzapine treatment was continued. Among 2500 patients in oral olanzapine 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).

Potential for Cognitive and Motor Impairment—Somnolence was a commonly reported, dose-related adverse event in premarketing trials (olanzapine 26% vs placebo 15%). Somnolence led to discontinuation in 0.4% (9/2500) of patients in the oral premarketing database.

Body Temperature Regulation—Use appropriate care when prescribing olanzapine for patients who will be experiencing conditions that may contribute to an elevation in core body temperature. Dysphagia—Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration pneumonia is a common cause of morbidity and mortality in patients with advanced Alzheimer'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 should be written for the smallest quantity of tablets consistent with good patient management.

<u>Use in Patients with Concomitant Illnesses</u>—Olanzapine should be used with caution in patients with clinically significant prostatic hypertrophy, narrow angle glaucoma, or a history of paralytic 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 than with placebo: falls, somnolence, peripheral edema, abnormal gait, urinary incontinence, lethargy, increased weight, asthenia, pyrexia, pneumonia, dry mouth, visual hallucinations. Discontinuation due to adverse events was significantly greater with olanzapine than placebo (13% vs 7%). Elderly patients with dementia-related psychosis treated with olanzapine are at an increased risk of death compared to placebo. Olanzapine is not approved for treatment of patients with dementia-related psychosis. If the prescriber elects to treat this patient population, vigilance should be exercised (see BOX WARNING and WARNINGS).

Because of the risk of orthostatic hypotension with olanzapine, use caution in cardiac patients (see Hemodynamic Effects).

<u>Information for Patients</u>—See full prescribing information for information to discuss with patients taking planzapine.

<u>Laboratory Tests</u>—Periodic assessment of transaminases is recommended in patients with significant hepatic disease.

<u>Drug</u> 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 (e.g., 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 inhibition of a single enzyme may appreciably alter olanzapine clearance. A dosage adjustment may need to be considered with specific drugs.

Activated charcoal (1 g) reduced the C_{max} 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. Carbamazepine (200 mg bid) causes an approximately 50% increase in the clearance of olanzapine. Higher daily doses of carbamazepine may cause an even greater increase in olanzapine clearance. Neither ethanol (45 mg/70 kg single dose) nor warfarin (20 mg single dose) had an effect on olanzapine pharmacokinetics. Fluoxetine at 60 mg (single or multiple doses) causes a small increase in the C_{max} of olanzapine and a small decrease in olanzapine clearance; however, the impact of this factor is small in comparison to the overall variability between individuals, and dose modification is not routinely recommended. Fluoxamine decreases the clearance of olanzapine; lower doses of olanzapine should be considered in patients receiving fluvoxamine concomitantly. In vitro data suggest that a clinically significant pharmacokinetic interaction between olanzapine and valproate is unlikely.

Olanzapine is unlikely to cause clinically important drug interactions mediated by the enzymes CYP1A2, CYP2C9, CYP2C19, CYP2D6, and CYP3A. Single doses of olanzapine did not affect the pharmacokinetics of imipramine/desipramine or warfarin. Multiple doses of olanzapine did not influence the kinetics of diazepam/N-desmethyldiazepam, lithium, ethanol, or biperiden. However, coadministration of either diazepam or ethanol potentiated the orthostatic hypotension observed with olanzapine. Multiple doses of olanzapine did not affect the pharmacokinetics of theophylline or its metabolites. Co-administration of intramuscular lorazepam and intramuscular olanzapine for injection added to the somnolence observed with either drug alone (see Hemodynamic 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 nortalities in males in the 30/20 mg/kg/d group. The incidence of mammary gland adenomas and adenocarcinomas was significantly increased in female mice and rats 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-fold 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). Diestrous was prolonged and estrous delayed at 0.6 times the MHDOD (mg/m² basis); therefore, olanzapine may produce a delay in ovulation.

<u>Pregnancy Category C</u>—There are no adequate and well-controlled studies in pregnant women. Olanzapine should be used in pregnancy only if the potential benefit justifies the potential risk to the fetus.



Start and stay with nonscheduled Rozerem— ZERO evidence of abuse or dependence



Clinical studies show no evidence of potential abuse, dependence, or withdrawal*

- First and only—nonscheduled prescription insomnia medication...not a controlled substance and approved for long-term use¹
- **First and only**—prescription insomnia medication that targets the normal sleep-wake cycle¹
- First and only—prescription insomnia medication with no evidence of abuse potential in clinical studies¹
- First and only—prescription insomnia medication that does not promote sleep by CNS depression¹
- Promote sleep with Rozerem—patients who took Rozerem fell asleep faster than those who took placebo¹
- One simple 8-mg dose¹

*Rozerem is not a controlled substance. A clinical abuse liability study showed no differences indicative of abuse potential between Rozerem and placebo at doses up to 20 times the recommended dose (N=14). Three 35-day insomnia studies showed no evidence of rebound insomnia or withdrawal symptoms with Rozerem compared to placebo (N=2082).¹²

Please visit www.rozerem.com

Rozerem is indicated for the treatment of insomnia characterized by difficulty with sleep onset. Rozerem can be prescribed for long-term use. Rozerem should not be used in patients with hypersensitivity to any components of the formulation, severe hepatic impairment, or in combination with fluvoxamine. Failure of insomnia to remit after a reasonable period of time should be medically evaluated, as this may be the result of an unrecognized underlying medical disorder. Hypnotics should be administered with caution to patients exhibiting signs and symptoms of depression. Rozerem has not been studied in patients with severe sleep apnea, severe COPD, or in children or adolescents. The effects in these populations are unknown. Avoid taking Rozerem with alcohol. Rozerem has been associated with decreased testosterone levels and increased prolactin levels. Health professionals should be mindful of any unexplained symptoms possibly associated with such changes in these hormone levels. Rozerem should not be taken with or immediately after a high-fat meal. Rozerem should be taken within 30 minutes before going to bed and activities confined to preparing for bed. The most common adverse events seen with Rozerem that had at least a 2% incidence difference from placebo were somnolence, dizziness, and fatigue.

Please see adjacent Brief Summary of Prescribing Information.



Proven for sleep. Nonscheduled for added safety.





Brief Summary of Prescribing Information 05-1114

ROZEREM™

INDICATIONS AND USAGE ROZEREM is indicated for the treatment of insomnia characterized by difficulty with sleep onset.

CONTRAINDICATIONS

ROZEREM is contraindicated in patients with a hypersensitivity to ramelteon or any components of the ROZEREM formulation.

Since sleep disturbances may be the presenting manifestation of a physical and/or psychiatric disorder, symptomatic treatment of insomnia should be initiated only after a careful evaluation of the patient. The failure of insomnia to remit after a reasonable period of treatment may indicate the presence of a primary psychiatric and/or medical illness that should be evaluated. Worsening of insomnia, or the emergence of new cognitive or behavioral abnormalities, may be the result of an unrecognized underlying psychiatric or physicial disorder and requires further evaluation of the patient. As with other hypnotics, exacerbation of insomnia and emergence of cognitive and behav-ioral abnormalities were seen with ROZEREM during the clinical development

ROZEREM should not be used by patients with severe hepatic impairment

ROZEREM should not be used in combination with fluvoxamine (see **PRE-CAUTIONS:** Drug Interactions).

A variety of cognitive and behavior changes have been reported to occur in association with the use of hypnotics. In primarily depressed patients, worsening of depression, including suicidal ideation, has been reported in association with the use of hypnotics.

Patients should avoid engaging in hazardous activities that require concentra-tion (such as operating a motor vehicle or heavy machinery) after taking ROZEREM.

After taking ROZEREM, patients should confine their activities to those necessary to prepare for bed.

PRECAUTIONS

General ROZEREM has not been studied in subjects with severe sleep apnea or NOZEMENT IS IN CHEEN STUDIED IN SUPPLIES WITH SEVERE SHEEP APIRE OF SEVERE COPP and is not recommended for use in those populations.

Patients should be advised to exercise caution if they consume alcohol in combination with ROZEREM.

Use in Adolescents and Children
ROZEREM has been associated with an effect on reproductive hormones in
adults, e.g. decreased testosterone levels and increased prolatein levels. It is
not known what effect chronic or even chronic intermittent use of ROZEREM
may have on the reproductive axis in developing humans (see Pediatric Use).

Information for Patients
Patients Patients Patients advised to take ROZEREM within 30 minutes prior to going to bed and should confine their activities to those necessary to prefor bed.

Patients should be advised to avoid engaging in hazardous activities (such as operating a motor vehicle or heavy machinery) after taking ROZEREM.

Patients should be advised that they should not take ROZEREM with or

immediately after a high fat meal. Patients should be advised to consult their health care provider if they experience worsening of insomnia or any new behavioral signs or symptoms of

Patients should consult their health care provider if they experience one of the following: cessation of menses or galactorrhea in females, decreased libido, or problems with fertility.

Laboratory Tests
No standard monitoring is required.

For patients presenting with unexplained amenorrhea, galactorrhea, decreased libido, or problems with fertility, assessment of prolactin levels and testosterone levels should be considered as appropriate.

Drug Interactions

ROZEREM has a highly variable inter-subject pharmacokinetic profile
(approximately 100% coefficient of variation in C_{max} and AUC). As noted
above, CYP1A2 is the major isozyme involved in the metabolism of
ROZEREM; the CYP2C subfamily and CYP3A4 isozymes are also involved
to a minor degree.

to a minor degree.
Effects of Other Drugs on ROZEREM Metabolism
Fluvoxamine (strong CYP1A2 Inhibitor): When fluvoxamine 100 mg twice
daily was administered for 3 days prior to single-dose co-administration of
ROZEREM 16m gan of fluvoxamine, the AUG_and for rameltoon increased
approximately 190-fold, and the C_{max} increased approximately 70-fold,
compared to ROZEREM administered alone. ROZEREM should not be used
in combination with fluvoxamine (See WARNINGS). Other less potent
CYP1A2 inhibitors have not been adequately studied. ROZEREM should be
administered with caution to patients taking less strong CYP1A2 inhibitors.
Rifampin (strong CYP enzyme inducer): Administration of rifampin 600 mg
once daily for 11 days resulted in a mean decrease of approximately 80%
(40% to 90%) in total exposure to ramelteon and metabolite M-II, (both
AUGG_aut and Cmi_m) after a single 32 mg dose of ROZEREM Efficacy may be
reduced when ROZEREM is used in combination with strong CYP enzyme
inducers such as rifampin.

inducers such as ritampin.

Ketoconzaole (strong CY9344 inhibitor): The AUC_{post} and C_{ross} of ramelteon increased by approximately 84% and 36%, respectively, when a single 16 mg does of ROZEREM was administered on the fourth day of ketoconazole good ong twice daily administration, compared to administration of ROZEREM alone. Similar increases were seen in M-II pharmacokinetic variables. ROZEREM should be administration in subjects taking strong CYP344 inhibitors such as ketoconazin in subjects taking strong CYP344 inhibitors such as ketoconazin.

Fluconazole (strong CYP2C9 inhibitor): The total and peak systemic exposure AUC_{o-int} and C_{max}) of ramelteon after a single 16 mg dose of ROZEREM was increased by approximately 150% when administered with fluconazole. Similar increases were also seen in M-II exposure. ROZEREM should be administered with caution in subjects taking strong CYP2C9 inhibitors such as fluconazole.

Interaction studies of concomitant administration of ROZEREM with fluoxe-tine (CYP206 inhibitor), omeprazole (CYP1A2 inducer/CYP2C19 inhibitor), theophylline (CYP1A2 substrate), and dextromethorphan (CYP206 substrate) did not produce clinically meaningful changes in either peak or total expo-sures to ramelteon or the M-II metabolite.

Sures to ratellection of ine W-II interactions.
Fiffects of ROZEREM on Metabolism of Other Drugs Concomitant administration of ROZEREM with omeprazole (CYP2C19 substrate), externethorphan (CYP2G6 substrate), midazolam (CYP3A4 substrate), theophylline (CYP1A2 substrate), diqoxin (p-glycoprotein substrate), and warfarin (CYP2G1 S)(CYP1A2 [IR] substrate) and warfarin (CYP2G1 S)(CYP1A2 [IR] substrate) did not produce clinically meaningful changes in peak and total exposures to these drugs.

Effect of Alcohol on Rozerem
Alcohol: With single-dose, daytime co-administration of ROZEREM 32 mg
and alcohol (0.6 g/kg), there were no clinically meaningful or statistically sig-

nificant effects on peak or total exposure to ROZEREM. However, an additive mindant eneus oil peak of total exposure to NOZENEM. Trovever, air adurus effect was seen on some measures of psychomotor performance (i.e., the Digit Symbol Substitution Test, the Psychomotor Vigilance Task Test, and a Visual Analog Scale to sedation) at some post-dose time polisis. No additive effect was seen on the Delayed Word Recognition Test. Because alcohol by itself impairs performance, and the intended effect of ROZEREM is to promote sleep, patients should be cautioned not to consume alcohol when using ROZEREM.

Drug/Laboratory Test Interactions
ROZEREM is not known to interfere with commonly used clinical laboratory tests. In addition, in vitro data indicate that ramelteon does not cause false-positive results for benzodiazepines, opiates, barbiturates, cocaine, cannabinoids, or amphetamines in two standard urine drug screening methods in vitro.

Carcinogenesis, Mutagenesis, and Impairment of Fertility

Carcinogenesis, wilcagenesis, and impairment or remity Carcinogenesis and tages of 2,03,00,300, 300, or 1000 mg/kg/day by oral gavage. Male mice exhibited a dose-related increase in the incidence of hepatic tumors at dose levels >100 mg/kg/day incidence of hepatic tumors at dose levels >100 mg/kg/day incideng hepatic adenoma, hepatic carcinoma, and hepatoblastoma. Fermale mice developed a dose-related increase in the incidence of hepatic adenomas at dose levels >00 mg/kg/day of mg/kg/day dose level. The no-effect level for hepatic tumors in male mice was 30 mg/kg/day (105 elimes and 3-times the therapertic exposure to ramelteon and the active metabolite M-II, respectively, at the maximum recommended human dose [MRHD] based on an area-under-the-curve [AUC] comparison). The no-effect level for hepatic tumors in female mice was 100 mg/kg/day (105 -11-mes and 15 -1-mes the therapeutic exposure to ramelteon and M-II, respectively, at the MRHD based on AUC).

to rameltoon and M-II, respectively, at the MRHD based on AUCs). In a two-year carcinogenicity study conducted in the Sprague-Dawley rat, male and female rats were administered ramelteon at doses of 0, 15, 60, 250 or 1000 mg/kg/day by oral gavage. Male rats exhibited a dose-related increase in the incidence of hepatic adenoma and benign Leydig cell tumors of the testis at dose levels ≥ 250 mg/kg/day and hepatic carcinoma at the 1000 mg/kg/day dose level. Female rats exhibited a dose-related increase in the incidence of hepatic adenoma at dose levels ≥ 60 mg/kg/day and hepatic carcinoma at the 1000 mg/kg/day dose level. The no-effect level for hepatic tumors and benign Leydig cell tumors in male rats was 50 mg/kg/day (1, 429-limes and 12-limes the therapeutic exposure to ramelteon and M-II, respectively, at the MRHD based on AUC). The no-effect level for hepatic tumors in female rats was 15 mg/kg/day (472-times and 16-times the therapeutic exposure to ramelteon and M-II, respectively, at the MRHD based on AUC).

The development of hepatic tumors in rodents following chronic treatment The development of hepatic tumors in rodents following chronic treatment with non-genotoxic compounds may be secondary to microsomal enzyme induction, a mechanism for tumor generation not thought to occur in humans. Leydig cell tumor development following treatment with non-genotoxic compounds in rodents has been linked to reductions in circulating testosterone levels with compensatory increases in luteinizing hormone release, which is a known proliferative stimulus to Leydig cells in the rat testis. Rat Leydig cells are more sensitive to the stimulatory effects of luteinizing by gone then human Leydia cells. In greenbraic tending can be developed and the reduction of the properties therefore the control of the contr testis. An at Eyrily deviate are intre-sensitive of the simulatory entects, ulterinizing hormone than human Leydig cells. In mechanistic studies conducted in the rat, daily rameltenon administration at 250 and 1000 mg/kg/day for 4 weeks was associated with a reduction in plasma testosterone levels. In the same study, luteinizing hormone levels were elevated over a 24 hour. period after the last ramelteon treatment; however, the durability of this luteinizing hormone finding and its support for the proposed mechanistic explanation was not clearly established.

explanation was not userally established at Although the rodent tumors observed following ramelteon treatment occurred at plasma levels of ramelteon and M-II in excess of mean clinical plasma con-centrations at the MRHD, the relevance of both rodent hepatic tumors and benign rat Leydig cell tumors to humans is not known.

Mutagenesis

Ramelteon was not genotoxic in the following: in vitro bacterial reverse mutarameterows as say, in into mammalian cell gene mutation assay using the mouse lymphoma TK* cell line; in involvin vitro uscheduled DNA synthesis assay in rath pedatocytes, and in in involvin vitro nucleus assays conducted in mouse and rat. Ramelteon was positive in the chromosomal aberration assay in chiene harmster lung cells in the presence of S9 metabolic activation. Reparts studies indicated that the concentration of the M-II metabolite formed by the rat liver S9 fraction used in the *in vitro* genetic toxicology studies described above, exceeded the concentration of ramelteon; therefore, the genotoxic potential of the M-II metabolite was also assessed in these studies.

Impairment of Fertility

Impairment of Fertility
Ramelteon was administered to male and female Sprague-Dawley rats in an initial fertility and early embryonic development study at dose levels of 6, 60, or 600 mg/kg/day. No effects on male or female mating or fertility were observed with a ramelteon dose up to 600 mg/kg/day (786-times higher than the MRHD on a mg/m² basis). Irregular estrus cycles, reduction in the number of implants, and reduction in the number of implants, and reduction in the number of implants, and reduction in the number of orpora lutea occurred at the 600 mg/kg/day dose level. Administration of ramelteon up to 600 mg/kg/day dose level. Administration of ramelteon up to 600 mg/kg/day to male rats for 7 weeks had no effect on sperm quality and when the treated male rats were mated with untreated female rats there was no effect on a female rats there was no effect on ramelteon at 20, 60 or 200 mg/kg/day for the same study duration, females demonstrated irregular estrus, cycles with doses ≥ 00 mg/kg/day, but no effects were seen on implantation or embryo viability. The no-effect dose for fertility employints was 20 mg/kg/day in temales (26-times the MRHD on a mg/m² basis) when considering all studies.

Pregnancy: Pregnancy Category C

on a migrin basis) when considering an studies.

Pregnancy: Pregnancy Category C
Ramelteon has been shown to be a developmental teratogen in the rat when given in doses 197 times higher than the maximum recommended human dose (MRHD) on a mg/m² basis. There are no adequate and well-controlled studies in pregnant women. Ramelteon should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

studies in pregnant women. Ramelteon should be used during pregnancy only if the potential benefit justifies the potential risk to the feture. The effects of ramelteon on embryo-fetal development were assessed in both the rat and rabbit. Pregnant rats were administered ramelteon by oral gavage at doses of 0. 10. 40, 150, or 600 mg/kg/day during gestation days 6-17, which is the period of organopenesis in this species. Evidence of maternal toxicity and fetal teratogenicity was observed at doses greater than or equal to 150 mg/kg/day. Maternal toxicity was chiefly characterized by decreased obdy weight and, at 600 mg/kg/day, ataxia and decreased spontaneous movement. At maternally toxic doses (150 mg/kg/day) ergater), the fetuses demonstrated visceral malformations consisting of diaphragmatic herria and minor anatomical variations of the skeleton (irregularly shaped scapula). At 600 mg/kg/day (reductions in fetal body weight and malformations including cysts on the external genitalia were additionally observed. The no-effect level for teratogenicity in this study was 40 mg/kg/day (1,892-times and 45-times higher than the therapeutic exposure to ramelteon and the active metabolite M-II, respectively, at the MHIP based on an area-under-the-curve (AUC) comparison). Pregnant rabbits were administered ramelteon by oral gavage at doses of 0.12, 60, or 300 mg/kg/day, no evidence of fetal effects or teratogenicity was associated with any dose level. The no-effect level for teratogenicity was, therefore, 300 mg/kg/day (11,862-times and 99-times

higher than the therapeutic exposure to ramelteon and M-II, respectively, at the MRHD based on AUC).

higher than the therapeutic exposure to ramelteon and M-II, respectively, at the MRHD based on AUC). The effects of ramelteon on pre- and post-natal development in the rat were studied by administration of ramelteon to the pregnant rat by oral gavage at doses of 0, 30, 100, or 300 mg/kg/day from day 6 of gestation through parturition to postnatal (location) day 21, at which time offspring were weaned. Maternal toxicity was noted at doses of 100 mg/kg/day or greater and consisted of reduced body weight during the post-wearing period was also noticed in the offspring of the groups given 100 mg/kg/day and higher. Offspring in the 300 mg/kg/day group demonstrated physical and developmental delays including delayed cruption of the lover incisors, a delayed acquisition of the righting reflex, and an alteration of emotional response. These delays are often observed in the presence of reduced offspring oby weight but may still be indicative of developmental delay. An apparent decrease in the viability of offspring in the 300 mg/kg/day group also showed evidence of diaphragmatic hernia, a finding observed in the embryo-fetted development study previously described. There were no effects on the reproductive capacity of offspring and the resulting progeny were not different from those of vehicle-treated offspring. The no-effect level for pre- and postnatal development in this study was 30 mg/kg/day (30-times higher than the MRHO on a mg/m² basis).

Labor and Delivery
The potential effects of ROZEREM on the duration of labor and/or delivery, for either the mother or the fetus, have not been studied. ROZEREM has no established use in labor and delivery.

Nursing Mothers

Ramelteon is secreted in to the milk of lactating rats. It is not known whether this drug is excreted in human milk. No clinical studies in nursing mothers have been performed. The use of ROZEREM in nursing mothers is not

Pediatric Use
Safety and effectiveness of ROZEREM in pediatric patients have not been
established. Further study is needed prior to determining that this product
may be used safely in pre-pubescent and pubescent patients.

Geriatric Use
A total of 654 subjects in double-blind, placebo-controlled, efficacy trials who A total of 654 subjects in double-billind, placebo-controlled, efficacy trials who received ROZEREM were at least 65 years of age; of these, 199 were 75 years of age or older. No overall differences in safety or efficacy were observed between elderly and younger adult subjects.

ADVERSE REACTIONS

Overview

The data described in this section reflect exposure to ROZEREM in 4251 subjects, including 346 exposed for 6 months or longer, and 473 subjects for one year.

Adverse Reactions Resulting in Discontinuation of Treatment

Auverse reactions resulting in Discontinuation of Treatment Five percent of the 3594 individual subjects exposed to ROZTREM in clinical studies discontinued treatment owing to an adverse event, compared with 2% of the 1370 subjects receiving placebo. The most frequent adverse events leading to discontinuation in subjects receiving ROZEREM were somnolence (0.8%), dizzness (0.5%), nausea (0.3%), fatigue (0.3%), headache (0.3%), and insommia (0.3%).

and insomnia (0.3%).

ROZEREM Most Commonly Observed Adverse Events in Phase 1-3 trials

The incidence of adverse events during the Phase 1 through 3 trials

(% placebo, n=1370; % ramelleon [8 mg], n=1250) were: headache NOS

(7%, 7%), somnolence (3%, 5%), fatigue (2%, 4%), dizziness (3%, 5%), nausea (2%, 3%), insomnia exacerbated (2%, 3%), upper respiratory tract
infection NOS (2%, 3%), distrea NOS (2%, 2%), mayliga (1%, 2%),
depression (1%, 2%), dysgeusia (1%, 2%), attrialgia (1%, 2%),
influenza (0. 1%), blood corticol decreased (0.1%)

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 clinical trials of other drugs, and may not reflect the rates observed in practice. The adverse reaction information from clinical trials

does, however, provide a basis for identifying the adverse events that appear to be related to drug use and for approximating rates.

DRUG ABUSE AND DEPENDENCE

DRUG ABUSE AND DEPENDENCE

DROW ABOSE AND PERFORMENCE ROZEREM Is not a controlled substance. Human Data: See the CLINICAL TRIALS section, Studies Pertinent to Safety Concerns for Sleep-Promoting Agents in the Complete Prescribing Information.

Animal Data. Ramelteon did not produce any signals from animal behavioral studies indicating that the drug produces rewarding effects. Monkeys did not self-administer ramelteon and the drug did not induce a conditioned place preference in rats. There was no generalization between ramelteon and midazolam. Ramelteon did not affect rotorod performance, an indicator of disruption of motor function, and it did not potentiate the ability of diazepam to interfere with rotorod performance.

to interfere with roution periodinals or in humans after chronic adminis-tration did not produce withdrawal signs. Ramelteon does not appear to produce physical dependence.

OVERDOSAGE

Signs and Symptoms No cases of ROZEREM overdose have been reported during clinical develop-

ROZEREM was administered in single doses up to 160 mg in an abuse liability trial. No safety or tolerability concerns were seen

ity trial. No safety or tolerability concerns were seen. Recommended Treatment General symptomatic and supportive measures should be used, along with immediate pastric lavage where appropriate. Intravenous fluids should be administered as needed. As in all cases of drug overdose, respiration, pulse, blood pressure, and other appropriate vital signs should be monitored, and general supportive measures employed.

Hemodialysis does not effectively reduce exposure to ROZEREM. Therefore, the use of dialysis in the treatment of overdosage is not appropriate.

The use of diarysis in the treatment of overdosage is not appropriate. Poison Control Center As with the management of all overdosage, the possibility of multiple drug ingestion should be considered. The physician may contact a poison control center for current information on the management of overdosage.

Rx only

Rx only
Manufactured by:
Takeda Pharmaceutical Company Limited
540-8645 Osaka, JAPAN

Manufactured in: Takeda Ireland Ltd. Kilruddery, County Wicklow, Republic of Ireland

Marketed by:
Takeda Pharmaceuticals America, Inc.
475 Half Day Road
Lincolnshire, IL 60069
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References: 1. Rozerem package insert, Takeda Pharmaceuticals America, Inc. 2. Johnson MW, Suess PE, Griffiths RR. Ramelteon: a novel hypnotic lacking abuse liability and sedative side effects. Arch Gen Psychiatry, In press.

A POWERFUL SSRI that's well tolerated





UP TO 90% of depressed patients present with symptoms of anxiety²

PROVEN EFFICACY for Major Depressive Disorder and Generalized Anxiety Disorder³



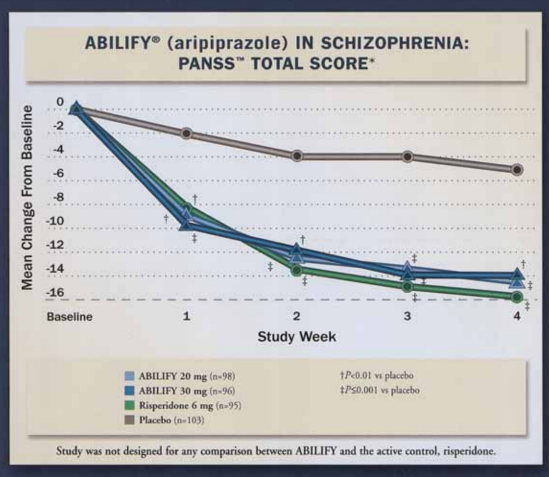
IMPORTANT SAFETY INFORMATION – Depression is a serious condition that can lead to suicidal thoughts and behavior. Antidepressants increased the risk of suicidal thinking and behavior (2% to 4%) in short-term studies of 9 antidepressant drugs in children and adolescents with major depressive disorder (MDD) and other psychiatric disorders. Patients started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior, especially at the beginning of therapy or at the time of dose changes. This risk may persist until significant remission occurs. Families and caregivers should be advised of the need for close observation and communication with the prescriber. Lexapro is not approved for use in pediatric patients.

Lexapro is contraindicated in patients taking monoamine oxidase inhibitors [MAOIs], pimozide [see DRUG INTERACTIONS – Pimozide and Celexa], or in patients with hypersensitivity to escitalopram oxalate. As with other SSRIs, caution is indicated in the coadministration of tricyclic antidepressants [TCAs] with Lexapro. As with other psychotropic drugs that interfere with serotonin reuptake, patients should be cautioned regarding the risk of bleeding associated with the concomitant use of Lexapro with NSAIDs, aspirin, or other drugs that affect coagulation. The most common adverse events with Lexapro versus placebo [approximately 5% or greater and approximately 2x placebo] were nausea, insomnia, ejaculation disorder, somnolence, increased sweating, fatigue, decreased libido, and anorgasmia.

References: 1. IMS National Prescription Audit. May 2005. 2. Sadock BJ, Sadock VA, Kaplan and Sadock's Synopsis of Psychiatry: Behavioral Sciences/Clinical Psychiatry: 9th ed. Philadelphia, Pa: Lippincott Williams & Wilkins; 2003:552. 3. LEXAPRO [package insert]. St Louis, Mo: Forest Pharmaceuticals, Inc.; 2005.

Please see brief summary of prescribing information for LEXAPRO on following page.

Controlling symptoms is one thing...



Data from a 4-week, double-blind, randomized, placebo-controlled trial (n=404) in hospitalized patients with acute schizophrenia or schizoaffective disorder. Mean baseline PANSS" Total Scores: ABILIFY 20 mg 94.4, ABILIFY 30 mg 92.6, risperidone 6 mg 94.9, placebo 95.7. Mean change from baseline: ABILIFY 20 mg -14.5, ABILIFY 30 mg -13.9, risperidone 6 mg -15.7, placebo -5.0.

Potkin et al. Arch Gen Psychiatry. 2003.

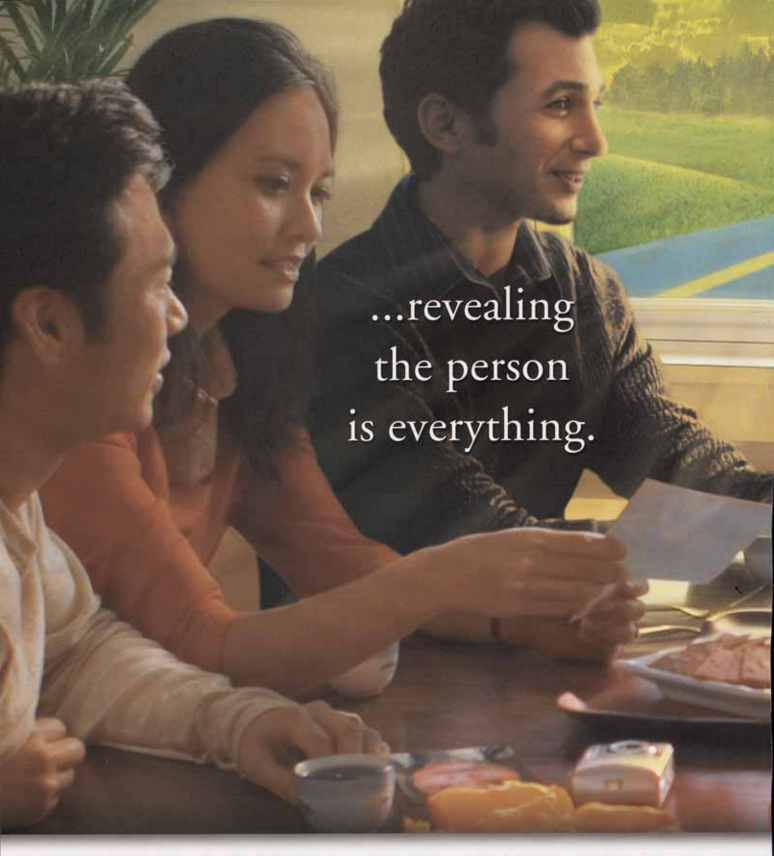
*Last observation carried forward (LOCF).

PANSS" (Positive and Negative Syndrome Scale) is a trademark of Multi-Health Systems, Inc.

ABILIFY is indicated for the treatment of schizophrenia.

A recommended starting and target dose of 10 or 15 mg/day is effective.

Please see IMPORTANT SAFETY INFORMATION, including **Bolded WARNING**, and INDICATIONS on page 4.



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. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular or infectious in nature.

ABILIFY is not approved for the treatment of patients with dementia-related psychosis.

Please see IMPORTANT SAFETY INFORMATION and INDICATIONS on page 4.





HELP ILLUMINATE THE PERSON WITHIN

IMPORTANT SAFETY INFORMATION and INDICATIONS for ABILIFY® (aripiprazole)

IMPORTANT SAFETY INFORMATION:

- 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. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular or infectious in nature. ABILIFY (aripiprazole) is not approved for the treatment of patients with dementia-related psychosis (see Boxed WARNING).
- ABILIFY is contraindicated in patients with a known hypersensitivity to the product.
- As with all antipsychotic medications, including ABILIFY, a rare condition referred to as neuroleptic malignant syndrome (NMS) has been reported. As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of tardive dyskinesia (TD).
- Cerebrovascular adverse events (eg, stroke, transient ischemic attack), including fatalities, have been reported at an increased incidence in clinical trials of elderly patients with dementia-related psychosis treated with ABILIFY, including a significant dose response relationship in a fixed-dose trial. ABILIFY is not approved for the treatment of patients with dementia-related psychosis.
- Hyperglycemia, including some serious cases ranging from ketoacidosis, hyperosmolar coma, or death, has been reported in patients treated with atypical antipsychotics. Patients on ABILIFY should be appropriately tested before and monitored during treatment.

ABILIFY may be associated with orthostatic hypotension and should be used with caution in patients with known cardiovascular disease, cerebrovascular disease, or conditions which would predispose them to hypotension.

As with other antipsychotic drugs, ABILIFY should be used with caution in patients with a history of seizures or with conditions that lower the seizure threshold.

Like other antipsychotics, ABILIFY may have the potential to impair judgment, thinking, or motor skills. Patients should not drive or operate hazardous machinery until they are certain ABILIFY does not affect them adversely.

Disruption of the body's ability to reduce core body temperature has been attributed to antipsychotics. Appropriate care is advised for patients who may exercise strenuously, be exposed to extreme heat, receive concomitant medication with anticholinergic activity, or be subject to dehydration.

As antipsychotics have been associated with esophageal dysmotility and aspiration, ABILIFY should be used cautiously in patients at risk for aspiration pneumonia.

As the possibility of a suicide attempt is inherent in psychotic illness and bipolar disorder, close supervision of high-risk patients should accompany drug therapy. Prescriptions for ABILIFY (aripiprazole) should be written for the smallest quantity consistent with good patient management to reduce the risk of overdose.

Physicians should determine if a patient is pregnant or intends to become pregnant while taking ABILIFY. Patients should be advised not to breast-feed while taking ABILIFY.

Patients should be advised to avoid alcohol while taking ABILIFY. Both CYP3A4 and CYP2D6 are responsible for ABILIFY metabolism. Agents that induce CYP3A4 (eg, carbamazepine) could cause an increase in ABILIFY clearance and lower blood levels. Inhibitors of CYP3A4 (eg, ketoconazole) or CYP2D6 (eg, quinidine, fluoxetine, or paroxetine) can inhibit ABILIFY elimination and cause increased blood levels.

Commonly observed adverse events reported with ABILIFY in 3-week bipolar mania trials at a ≥5% incidence for ABILIFY and at a rate at least twice the rate of placebo include, respectively, akathisia (15% vs 4%), constipation (13% vs 6%), and accidental injury (6% vs 3%).

Treatment-emergent adverse events reported with ABILIFY in short-term trials at an incidence ≥10% and greater than placebo, respectively, include headache (31% vs 26%), agitation (25% vs 24%), anxiety (20% vs 17%), insomnia (20% vs 15%), nausea (16% vs 12%), dyspepsia (15% vs 13%), somnolence (12% vs 8%), akathisia (12% vs 5%), lightheadedness (11% vs 8%), vomiting (11% vs 6%), and constipation (11% vs 7%).

The adverse events reported in a 26-week, double-blind schizophrenia trial comparing ABILIFY and placebo were generally consistent with those reported in the short-term, placebo-controlled schizophrenia trials, except for a higher incidence of tremor: 9% for ABILIFY vs 1% for placebo.

INDICATIONS: ABILIFY is indicated for the treatment of:

- Schizophrenia, including maintaining stability in patients who had been symptomatically stable on other antipsychotic medications for periods of 3 months or longer and observed for relapse during a period of up to 26 weeks*
- Acute manic and mixed episodes associated with Bipolar I Disorder
- Maintaining efficacy in patients with Bipolar I Disorder with a recent manic or mixed episode who had been previously stabilized and then maintained for at least 6 weeks*
- *Physicians who elect to use ABILIFY for extended periods should periodically re-evaluate the long-term usefulness of the drug for the individual patient.

ABILIFY is taken once daily with or without food.

Please see BRIEF SUMMARY of FULL PRESCRIBING INFORMATION, including **Boxed WARNING**, on the following pages.

New ABILIFY® DISCMELT (aripiprazole)

Orally Disintegrating Tablets



Similar efficacy and safety to ABILIFY tablets No liquid needed Rapidly disintegrates Convenient delivery of an effective dose



HELP ILLUMINATE THE PERSON

ABILIFY® (aripiprazole) Tablets ABILIFY® DISCMELT™ (aripiprazole) **Orally Disintegrating Tablets**

ABILIFY® (aripiprazole) Oral Solution

Brief Summary of Prescribing Information. For complete prescribing information please consult official package circular

WARNING

Increased Mortality in Elderly Patients with Dementia-Related Psychosis

Increased Mortality in Eiderly 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 (e.g., heart failure, sudden death) or infectious (e.g., pneumonia) in nature. ABILIFY (aripiprazole) is not approved for the treatment of patients with dementia-related psychosis.

CONTRAINDICATIONS

ABILIFY (aripiprazole) is contraindicated in patients with a known hypersensitivity to the product.

WARNINGS

WARNINGS
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. ABILIFY is not approved for the treatment of patients
with dementia-related psychosis (see Boxed WARNING).

Neuroleptic Malignant Syndrome (NMS)

Neuroleptic Malignant Syndrome (NMS)

A potentially fatal symptom complex sometimes referred to as Neuroleptic Malignant Syndrome (NMS) has been reported in association with administration of antipsychotic drugs, including aripiprazole. Two possible cases of NMS occurred during aripiprazole treatment in the premarketing worldwide clinical database. Clinical manifestations of NMS are hyperpyrexia, muscle rigidity, aftered mentistatius, and evidence of autonomic instability (irregular pulse or blood pressure, tachycardia, diaphoresis, and cardiac dysrhythmia). Additional signs may include elevated creatine phosphokinase, myoglobinuria (rhabdomyolysis), and acute renal failure.

The diannostic evaluation of nations with this syndrome is complicated. In article 1.

(rhabdomyolysis), and acute renal failure.

The diagnostic evaluation of patients with this syndrome is complicated. In arriving at a diagnosis, it is important to exclude cases where the clinical presentation includes both serious medical illness (e.g., pneumonia, systemic infection, etc) and untreated or inadequately treated extrapyramidal signs and symptoms (EPS). Other important considerations in the differential diagnosis include central artitcholinergic toxicity, heat stroke, drug fever, and primary central nervous system pathology.

The management of NMS should include: 1) immediate discontinuation of antipsychotic drugs and other drugs not essential to concurrent therapy; 2) intensive symptomatic treatment and medical monitoring; and 3) treatment of any concomitant serious medical problems for which specific treatments are available. There is no general agreement about specific pharmacological treatment regimens for uncomplicated NMS. If a patient requires anotipsychotic drug treatment after recovery from NMS, the potential reintroduction of drug therapy should be carefully considered. The patient should be carefully monitored, since recurrences of NMS have been reported.

Tardive Dyskinesia

Tardive Dyskinesia

A syndrome of potentially irreversible, involuntary, dyskinetic movements may develop in patients treated with antipsychotic drugs. Although the prevalence of the syndrome appears to be highest among the elderly, especially elderly women, it is impossible to rely upon prevalence estimates to predict, at the inception of antipsychotic treatment, which patients are likely to develop the syndrome. Whether antipsychotic drug products differ in their potential to cause tardive dyskinesia is unknown.

The risk of developing tardive dyskinesia and the likelihood that it will become irreversible are believed to

increase as the duration of treatment and the total cumulative dose of antipsychotic drugs administered to the patient increase. However, the syndrome can develop, although much less commonly, after relatively brief treatment periods at low doses.

treatment periods at low doses.

There is no known treatment for established cases of tardive dyskinesia, although the syndrome may remit, partially or completely, if antipsychotic treatment is withdrawn. Antipsychotic treatment, itself, however, may suppress (or partially suppress) the signs and symptoms of the syndrome and, thereby, may possibly mask the underlying process. The effect that symptomatic suppression has upon the long-term curse of the syndrome is unknown.

Given these considerations, ABILIFY should be prescribed in a manner that is most likely to minimize the occurrence of tardive dyskinesia. Chronic antipsychotic treatment should generally be reserved for patients who suffer from a chronic illness that (1) is known to respond to antipsychotic drugs, and (2) for whom alternative, equally effective, but potentially less harmful treatments are not available or appropriate in patients who do require chronic treatment, the smallest dose and the shortest duration of treatment producing a satisfactory clinical response should be sought. The need for continued treatment should be reassessed periodically.

periodically. periodically.

If signs and symptoms of tardive dyskinesia appear in a patient on ABILIFY, drug discontinuation should be considered. However, some patients may require treatment with ABILIFY despite the presence of the syndrome.

considered. However, some patients may require treatment with ABILIFY despite the presence of the syndrome.
Cerebrovascular Adverse Events, Including Stroke, in Elderly Patients with Dementia-Related Psychosis In placebo-controlled clinical studies (two flexible dose and one fixed dose study) of dementia-related psychosis, there was an increased incidence of cerebrovascular adverse events (e.g., stroke, transient ischemic attack), including fatalities, in aripiprazole-treated patients (mean age: 84 years; range: 78-88 years). In the fixed-dose study, there was a statistically significant dose response relationship for cerebrovascular adverse events in patients treated with aripiprazole. Aripiprazole is not approved for the treatment of patients with dementialed psychosis. (See also Boxed WaRNING, WaRNINGS: Increased Mortality in Elderly Patients with Dementia-Related Psychosis, and PRECAUTIONS: Use in Patients with Concomitant Illness: Safety Experience in Elderly Patients with Psychosis Associated with Alzheimer's Disease.)

Hyperglycemia and Diabetes Mellitus

Hyperglycemia, in some cases extreme and associated with ketoacidosis or hyperosmolar coma or death, has been reported in patients treated with atypical antipsychotics. There have been few reports of hyperglycemia in patients treated with ABILIFY, although fewer patients have been treated with ABILIFY, it is not known if this more limited experience is the sole reason for the paucity of such reports. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes mellitus in patients with schizophrenia and the increasing incidence of diabetes mellitus in the general population. Given these confounders, the relationship between atypical antipsychotic use and hyperglycemia-related adverse events is not completely understood. However, epidemiological studies which did not include ABILIFY suggest an increased risk of treatment-emergent hyperglycemia-related adverse events in patients treated with the atypical antipsychotics included in these studies. Because ABILIFY was not marketed at the time these studies were performed, it is not known if ABILIFY is associated with this increased risk. Precise risk estimates for hyperglycemia-related adverse events in patients treated with applical

marketed at the time these studies were performed, it is not known if ABILIFY is associated with this increased risk. Precise risk estimates for hyperglycemia-related adverse events in patients treated with atypical antipsychotics are not available.

Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics should be monitored regularly for worsening of glucose control. Patients with risk factors for diabetes mellitus (e.g., obesity, family history of diabetes) who are starting treatment with atypical antipsychotics should undergo fasting blood glucose testing at the beginning of treatment and periodically during treatment. Any patient treated with atypical antipsychotics should be monitored for symptoms of hyperglycemia including polydipsia, polypria, polyphagia, and weakness. Patients who develop symptoms of hyperglycemia during treatment with atypical antipsychotics should undergo fasting blood glucose testing. It nose cases, hyperglycemia has resolved when the atypical antipsychotic was discontinued; however, some patients required continuation of anti-diabetic treatment despite discontinuation of the suspect drug.

PRECAUTIONS

General

Orthostatic Hypotension

Aripiprazole may be associated with orthostatic hypotension, perhaps due to its α_1 -adrenergic receptor antagonism. The incidence of orthostatic hypotension-associated events from five short-term, placebo-

controlled trials in schizophrenia (n=926) on ABILIFY (aripiprazole) included: orthostatic hypotension (placebo 1%, aripiprazole 1.9%), orthostatic lightheadedness (placebo 1%, aripiprazole 0.9%), and syncope (placebo 1%, aripiprazole 0.6%). The incidence of orthostatic hypotension-associated events from short-term, placebo-controlled trials in bipolar mania (n=597) on ABILIFY included: orthostatic hypotension (beabo 0%, aripiprazole 0.7%), orthostatic lightheadedness (placebo 0.5%, aripiprazole 0.5%), and syncope (placebo 0.9%, aripiprazole 0.5%). The incidence of a significant orthostatic change in blood pressure (defined as a decrease of at least 30 mmHg in systolic blood pressure when changing from a supine to standling position) for aripiprazole was not statistically different from placebo (in schizophrenia: 14% among aripiprazole-treated patients and 12% among placebo-treated patients and in bipolar mania: 3% among aripiprazole-treated patients and 2% among placebo-treated patients.

placebo-treated patients).

placebo-treated patients).

Aripiprazio e should be used with caution in patients with known cardiovascular disease (history of myocardial infarction or ischemic heart disease, heart failure or conduction abnormalities), cerebrovascular disease, or conditions which would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications).

RONLY

Seizure
Seizures occurred in 0.1% (1/926) of aripiprazole-treated patients with schizophrenia in short-term, placebo-controlled trials. In short-term, placebo-controlled clinical trials of patients with bipolar mania, 0.3% (2/597) of aripiprazole-treated patients and 0.2% (1/436) of placebo-treated patients experienced seizures. As with other antipsychotic drugs, aripiprazole should be used cautiously in patients with a history of seizures or with conditions that lower the seizure threshold, e.g., Alzheimer's dementia. Conditions that lower the seizure threshold may be more prevalent in a population of 65 years or older.

threshold may be more prevalent in a population of 65 years of older.

Potential for Cognitive and Motor Impairment in short-term, placebo-controlled trials of schizophrenia, somnolence was reported in 11% of patients on ABILIFY compared to 8% of patients on of patients on controlled trials of schizophrenia, somnolence led to discontinuation in 0.1% (1/926) of patients with schizophrenia on ABILIFY in short-term, placebo-controlled trials of bipolar mania, somnolence was reported in 14% of patients on ABILIFY compared to 7% patients on Jacebo, but did not lead to discontinuation of any patients with bipolar mania. Despite the relatively modest increased incidence of somnolence compared to placebo, ABILIFY, like other antipsychotics, may have the potential to impair judgment, thinking, or motor skills. Patients should be cautioned about operating hazardous machinery, including automobiles, until they are reasonably certain that therapy with ABILIFY does not affect them adversely.

Rody Temperature Benutation

Body Temperature Regulation

Disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents.
Appropriate care is advised when prescribing arripiprazole for patients who will be experiencing conditions which may contribute to an elevation in core body temperature, e.g., exercising strenuously, exposure to extreme heat, receiving concomitant medication with anticholinergic activity, or being subject to dehydration.

Dysphagia

Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration
preumonia is a common cause of morbidity and mortality in elderly patients, in particular those with
advanced Alzheimer's dementia. Aripiprazole and other antipsychotic drugs should be used cautiously in
patients at risk for aspiration pneumonia (see PRECAUTIONS: Use In Patients with Concomitant Illness).

Suicide

The possibility of a suicide attempt is inherent in psychotic illnesses and bipolar disorder, and close supervision of high-risk patients should accompany drug therapy. Prescriptions for ABILIFY should be written for the smallest quantity consistent with good patient management in order to reduce the risk of overdose.

Use in Patients with Concomitant Iliness
Clinical experience with ABILIFY in patients with certain concomitant systemic illnesses (see CLINICAL PHARMACOLOGY: Special Populations: Renal Impairment and Hepatic Impairment in Full Prescribing

Information) is limited.

ABILIFY has not been evaluated or used to any appreciable extent in patients with a recent history of myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from

myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from premarketing clinical studies.

Safety Experience in Elderly Patients with Psychosis Associated with Alzheimer's Disease: In three, 10-week, placebo-controlled studies of aripiprazole in elderly patients with psychosis associated with Alzheimer's disease (n=938; mean age: 82.4 years; range: 56-99 years), the treatment-emergent adverse events that were reported at an incidence of ≥3% and aripiprazole incidence at least twice that for placebo were asthenia (placebo 3%, aripiprazole 8%), somnolence (placebo 3%, aripiprazole 9%), urinary incontinence (placebo 1%, aripiprazole 5%), excessive salivation (placebo 3%, aripiprazole 4%), and lightheadedness (placebo 1%, aripiprazole 4%).

The safety and efficacy of ABILIFY in the treatment of patients with psychosis associated with dementia have not been established. If the prescriber elects to treat such patients with ABILIFY, vigilance should be exercised, particularly for the emergence of difficulty swallowing or excessive somnolence, which could predispose to accidental injury or aspiration. (See also Boxed WARMING, WARMINGS: Increased Mortality in Elderty Patients with Dementia-Related Psychosis, and Cerebrovascular Adverse Events, Including Stroke, in Elderly Patients with Dementia-Related Psychosis.)

Information for Patients

Physicians are advised to discuss the following issues with patients for whom they prescribe ABILIFY:

Interference with Cognitive and Motor Performance: Because aripiprazole may have the potential to impair judgment, thinking, or motor skills, patients should be cautioned about operating hazardous machinery, including automobiles, until they are reasonably certain that aripiprazole therapy does not affect them adversely

Pregnancy: Patients should be advised to notify their physician if they become pregnant or intend to become pregnant during therapy with ABILIFY.

Nursing: Patients should be advised not to breast-feed an infant if they are taking ABILIFY.

Concomitant Medication: Patients should be advised to inform their physicians if they are taking, or plan to take, any prescription or over-the-counter drugs, since there is a potential for interactions.

Alcohol: Patients should be advised to avoid alcohol while taking ABILIFY.

Heat Exposure and Dehydration: Patients should be advised regarding appropriate care in avoiding overheating Sugar Content: Patients should be advised that each mL of ABILIFY oral solution contains 400 mg of sucrose and 200 mg of fructose.

Phenylketonurics: Phenylalanine is a component of aspartame. Each ABILIFY DISCMELT orally disintegrating tablet contains the following amounts: 10 mg - 1.12 mg phenylalanine and 15 mg - 1.68 mg phenylalanine.

Drug-Drug Interactions

Given the primary CNS effects of aripiprazole, caution should be used when ABILIFY is taken in combination with other centrally acting drugs and alcohol. Due to its α_1 -adrenergic receptor antagonism, aripiprazole has the potential to enhance the effect of certain antihypertensive agents.

Potential for Other Drugs to Affect ABILITY
Ariphtracole is not a substrate of CYP1A1, CYP1A2, CYP2A6, CYP2B6, CYP2C8, CYP2C9, CYP2C19, or CYP2E1

Potential for Other Drugs to Artect ABILIPY
Aripiprazole is not a substrate of CYP1A1, CYP1A2, CYP2A6, CYP2B6, CYP2C8, CYP2C9, CYP2C19, or CYP2E1
enzymes. Aripiprazole also does not undergo direct glucuronidation. This suggests that an interaction of
aripiprazole with inhibitors or inducers of these enzymes, or other factors, like smoking, is unlikely.

Both CYP3A4 and CYP2D6 are responsible for aripiprazole metabolism. Agents that induce CYP3A4 (e.g.,
carbamazepine) could cause an increase in aripiprazole clearance and lower blood levels. Inhibitors of
CYP3A4 (e.g., ketoconazole) or CYP2D6 (e.g., quinifine, fluoxettine, or paroxettine) can inhibit aripiprazole
elimination and cause increased blood levels.

Ketoconazole: Coadministration of ketoconazole (200 mg/day for 14 days) with a 15-mg single dose of
aripiprazole increased the AUC of aripiprazole and its active metabolite by 63% and 77%, respectively. The
effect of a higher ketoconazole dose (400 mg/day) has not been studied. When concomitant administration of
ketoconazole with aripiprazole occurs, aripiprazole dose should be reduced to one-half of its normal dose.
Other strong inhibitors of CYP3A4 (firaconazole) would be expected to have similar effects and need similar
dose reductions; weaker inhibitors (erythromytin, grapefruit juice) have not been studied. When CYP3A4
inhibitor is withdrawn from the combination therapy, aripiprazole dose should then be increased.

Quinidine: Coadministration of a 10-mg single dose of aripiprazole by 112% but decreased the AUC of its
active metabolite, dehydro-aripiprazole, by 35%. Aripiprazole dose should be reduced to one-half of its
normal dose when concomitant administration of quinidine with aripiprazole occurs. Other significant
inhibitors of CYP2D6, such as fluoxetine or paroxetine, would be expected to have similar effects and,
therefore, should be accompanied by similar dose reductions. When the CYP2D6 inhibitor is withdrawn from
the combination therapy, aripiprazole dose should then be increased.

Carbamazepine: Coadministration of carbamazepine (200 mg BID), a potent CYP3A4 inducer, with aripiprazole (30 mg QD) resulted in an approximate 70% decrease in $C_{\rm max}$ and AUC values of both aripiprazole and its active metabolite, dehydro-aripiprazole. When carbamazepine is added to aripiprazole therapy, aripiprazole dose should be doubled. Additional dose increases should be based on clinical evaluation. When carbamazepine is withdrawn from the combination therapy, aripiprazole dose should then

No clinically significant effect of famotidine, valproate, or lithium was seen on the pharmacokinetics of aripiprazole (see CLINICAL PHARMACOLOGY: Drug-Drug Interactions in Full Prescribing Information).

aripiprazole (see CLINICAL PHARMACOLOGY: Drug-Drug interactions in Full Prescribing Information).
Potential for ABILIFY (aripiprazole) to Affect Other Drugs
Aripiprazole is unlikely to cause clinically important pharmacokinetic interactions with drugs metabolized by cytochrome P450 enzymes. In in vivo studies, 10- to 30-my/day doses of aripiprazole had no significant effect on metabolism by CYP206 (dextromethorphan), CYP209 (warfarin), CYP2C19 (omeprazole, warfarin), and CYP3A4 (dextromethorphan) substrates. Additionally, aripiprazole and dehydro-aripiprazole did not show potential for altering CYP1A2-mediated metabolism in vitro (see CLINICAL PHARMACOLOGY: Drug-Drug Interactions in Full Prescribing Information).

**Alcohol:* There was no significant difference between aripiprazole coadministered with ethanol on performance of gross motor skills or stimulus response in healthy subjects. As with most psychoactive medications, patients should be advised to avoid alcohol while taking ABILIFY.

Carcinogenesis, Mutagenesis, Impairment of Fertility

Lifetime carcinogenicity studies were conducted in ICR mice and in Sprague-Dawley (SD) and F344 rats. Lifetime carcinogenicity studies were conducted in ICR mice and in Sprague-Dawley (SID) and F344 rats. Arpiprazole was administered for 2 years in the diet at doses of 1, 3, 10, and 30 mg/kg/day to Temice and 1, 3, and 10 mg/kg/day to F344 rats (0.2 to 5 and 0.3 to 3 times the maximum recommended human dose [MRHD] based on mg/m², respectively). In addition, SD rats were dosed orally for 2 years at 10, 20, 40, and 60 mg/kg/day (3 to 19 times the MRHD based on mg/m²). Aripiprazole did not induce tumors in male mice or rats. In female mice, the incidences of pituitary gland adenomas and mammary gland adenocarcinomas and adenocarcinomas were increased at dietary doses of 3 to 30 m/kg/day (0.1 to 0.9 times human exposure at MRHD based on AUC and 0.5 to 5 times the MRHD based on mg/m²). In female rats, the incidence of mammary gland fibroadenomas was increased at a dietary dose of 10 mg/kg/day (0.1 times human exposure at MRHD based on AUC and 3 times the MRHD based on mg/m²); and the incidences of adrenocortical adenomas/carcinomas were increased at an oral dose of 6 mg/kg/day (1.1 times human exposure at MRHD based on mg/m²) and the incidences of adrenocortical carcinomas and combined adrenocortical adenomas/carcinomas were increased at an oral dose of 6 mg/kg/day (0.1 to 10 mg/m²).

carcinomas and combined adrenocortical adenomas/carcinomas were increased at an oral dose of 60 my/kg/dyd (14 times human exposure at MRHD based on AUC and 19 times the MRHD based on my/m²). Proliferative changes in the pituitary and mammary gland of rodents have been observed following chronic administration of other antipsychotic agents and are considered prolactin-mediated. Serum prolactin was not measured in the arripprazole carcinogenicity studies. However, increases in serum prolactin was not observed in female mice in a 13-week dietary study at the doses associated with mammary gland and pituitary tumors. Serum prolactin was not increased in female rats in 4- and 13-week dietary studies at the dose associated with mammary gland tumors. The relevance for human risk of the findings of prolactin-mediated endocrine tumors in rodents is unknown.

Mutagenesis
The mutagenic potential of aripiprazole was tested in the *in vitro* bacterial reverse-mutation assay, the *in vitro* bacterial DNA repair assay, the *in vitro* forward gene mutation assay in mouse lymphoma cells, the *in vitro* chromosomal aberration assay in Chinese harmster lung (CHL) cells, the *in vitro* micronucleus assay in mice, and the unscheduled DNA synthesis assay in rats. Aripiprazole and a metabolite (2,3-DCPP) were clastogenic in the *in vitro* chromosomal aberration assay in CHL cells with and without metabolic activation. The metabolite, 2,3-DCPP, produced increases in numerical aberrations in the *in vitro* assay in CHL cells in the absence of metabolic activation. A positive response was obtained in the *in vivo* micronucleus assay in mice; however, the response was shown to be due to a mechanism not considered relevant to humans

Impairment of Fertility

Impaliment of Fertility
Female rats were treated with oral doses of 2, 6, and 20 mg/kg/day (0.6, 2, and 6 times the maximum recommended human dose [MRHD] on a mg/m² basis) of aripiprazole from 2 weeks prior to mating through day 7 of gestation. Estrus cycle irregularities and increased corpora lutea were seen at all doses, but no impairment of fertility was seen. Increased pre-implantation loss was seen at 6 and 20 mg/kg, and decreased fetal weight was seen at 20 mg/kg.

Male rats were treated with oral doses of 20, 40, and 60 mg/kg/day (6, 13, and 19 times the MRHD on a mg/m² basis) of aripiprazole from 9 weeks prior to mating through mating. Disturbances in spermatogenesis were seen at 60 mg/kg, and prostate atrophy was seen at 40 and 60 mg/kg, but no impairment of fertility was seen.

was seen.

Pregnancy Pregnancy Category C

in animal studies, aripiprazole demonstrated developmental toxicity, including possible teratogenic effects in rats and rabbits.

in animal studies, airpiprazole demonstrated developmental toxicity, including possible teratogenic effects in rats and rabbits.

Pregnant rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the maximum recommended human dose [MRHD] on a mg/m² basis) of aripiprazole during the period of organogenesis. Gestation was slightly prolonged at 30 mg/kg. Treatment caused a slight delay in fetal development, as evidenced by decreased fetal weight (30 mg/kg), undescended testes (30 mg/kg), and delayed skeletal ossification (10 and 30 mg/kg). There were no adverse effects on embryofetal or pup survivals. Delivered offspring had decreased bodyweights (10 and 30 mg/kg), and increased incidences of hepatodiaphragmatic nocules and diaphragmatic hernia at 30 mg/kg (the other dose groups were not examined for these findings). (A low incidence of diaphragmatic hernia was also seen in the fetuses exposed to 30 mg/kg.) Postnatally, delayed vaginal opening was seen at 10 and 30 mg/kg and impaired reproductive performance (decreased fertillity rate, corpora lutea, implants, and live fetuses, and increased post-implantation loss, likely mediated through effects on female offspring) was seen at 30 mg/kg, some maternal toxicity was seen at 30 mg/kg, however, there was no evidence to suggest that these developmental effects were secondary to maternal toxicity. Pregnant rabbits were treated with oral doses of 10, 30, and 100 mg/kg/day (2, 3, and 11 times human exposure at MRHD based on AUC and 6, 19, and 65 times the MRHD based on mg/m²) of aripiprazole during the period of organogenesis. Decreased maternal food consumption and increased fetal weight (30 and 100 mg/kg). Recreased fetal weight (30 and 100 mg/kg). Treatment caused increased maternal food consumption and increased fetal weight (30 and 100 mg/kg). Treatment caused increased fetal mortality (100 mg/kg), decreased fetal weight (30 and 100 mg/kg). In a study in which rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times th

There are no adequate and well-controlled studies in pregnant woman it is not known whether aripiprazole can cause fetal harm when administered to a pregnant woman or can affect reproductive capacity. Aripiprazole should be used during pregnancy only if the potential benefit outweighs the potential risk to the fetus.

Labor and Delivery

The effect of aripiprazole on labor and delivery in humans is unknown.

Nursing Mothers

Aripiprazole was excreted in milk of rats during lactation. It is not known whether aripiprazole or its metabolites are excreted in human milk, It is recommended that women receiving aripiprazole should not

Safety and effectiveness in pediatric and adolescent patients have not been established.

Geriatric Use

Of the 7951 patients treated with aripiprazole in premarketing clinical trials, 991 (12%) were ≥65 years old and 789 (10%) were ≥75 years old. The majority (88%) of the 991 patients were diagnosed with dementia of the Alzheimer's type.

Placebo-controlled studies of aripiprazole in schizophrenia or bipolar mania did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. There was no effect of age on the pharmacokinetics of a single 15-mg dose of aripiprazole. Aripiprazole clearance was decreased by 20% in elderly subjects (≥65 years) compared to younger adult subjects (18 to 64 years), but there was no detectable effect of age in the population pharmacokinetic analysis in schizophrenia patients.

scnrzopnrenia patients.

Studies of elderly patients with psychosis associated with Alzheimer's disease have suggested that there may be a different tolerability profile in this population compared to younger patients with schizophrenia (see Boxed WARNING; WARNINGS: Increased Mortality in Elderly Patients with Dementia-Related Psychosis; Cerebrovascular Adverse Events, Including Stroke, in Elderly Patients with Dementia-

Related Psychosis: and PRECAUTIONS: Use in Patients with Concomitant Illness). The safety and efficacy of ABILIFY (appiprazole) in the treatment of patients with psychosis associated with Alzheimer's disease has not been established. If the prescriber elects to treat such patients with ABILIFY, vigilance should be exercised.

ADVERSE REACTIONS

ADVERSE REACTIONS
Aripiprazole has been evaluated for safety in 7951 patients who participated in multiple-dose, premarketing trials in schizophrenia, bipolar mania, and dementia of the Alzheimer's type, and who had approximately 5235 patient-years of exposure. A total of 2280 aripiprazole-treated patients were treated for at least 180 days and 1558 aripiprazole-treated patients had at least 1 year of exposure.

The conditions and duration of treatment with aripiprazole included (in overlapping categories) double-blind, comparative and noncomparative open-label studies, inpatient and outpatient studies, fixed- and

blind, comparative and noncomparative open-label studies, inpatient and outpatient studies, liked-and flexible-dose studies, and short- and longer-term exposure.

Adverse events during exposure were obtained by collecting volunteered adverse events, as well as results of physical examinations, vital signs, weights, laboratory analyses, and ECG. Adverse experiences were recorded by clinical investigators using terminology of their own choosing. In the tables and tabulations that follow, modified COSTART dictionary terminology has been used initially to classify reported adverse events into a smaller number of standardized event categories, in order to provide a meaningful estimate of

events into a smaller number of standardized event categories, in order to provide a meaningful estimate of the proportion of individuals experiencing adverse events.

The stated frequencies of adverse events represent the proportion of individuals who experienced at least once, a treatment-mergent adverse event of the type listed. An event was considered treatment emergent if it occurred for the first time or worsened while receiving therapy following baseline evaluation. There was no attempt to use investigator causality assessments; ie, all reported events are included.

The prescriber should be aware that the figures in the tables and tabulations cannot be used to predict the incidence of side effects in the course of usual medical practice where patient characteristics and other factors differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatment, uses, and investigators. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of dring and pondrug factors to the adverse event incidence in the population studied. contribution of drug and nondrug factors to the adverse event incidence in the population studied.

Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Schizophrenia The following findings are based on a pool of five placebo-controlled trials (four 4-week and one 6-week) in which aripiprazole was administered in doses ranging from 2 to 30 mg/day.

Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials Overall, there was no difference in the incidence of discontinuation due to adverse events between aripiprazole-treated (7%) and placebo-treated (9%) patients. The types of adverse events that led to discontinuation were similar between the aripiprazole- and placebo-treated patients.

Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Ma The following findings are based on a pool of 3-week, placebo-controlled, bipolar mania trials in which aripiprazole was administered at doses of 15 or 30 mg/day.

Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials Overall, in patients with bipclar mania, there was no difference in the incidence of discontinuation due to adverse events between aripiprazole-treated (11%) and placebo-treated (9%) patients. The types of adverse events that led to discontinuation were similar between the aripiprazole and placebo-treated patients

Commonly Observed Adverse Events in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Mania Commonly observed adverse events associated with the use of aripiprazole in patients with bipolar mania (incidence of 5% or greater and aripiprazole incidence at least twice that for placebo) are shown in Table 1. There were no adverse events in the short-term trials of schizophrenia that met these criteria.

Table 1: Commonly Observed Adverse Events in Short-Term,
Placebo-Controlled Trials of Patients with Bipolar Mania

	Percentage of Patien	Percentage of Patients Reporting Event	
Adverse Event	Aripiprazole (n=597)	Placebo (n=436)	
Accidental Injury	6	3	
Constipation	13	6	
Akathisia	15	4	

Adverse Events Occurring at an Incidence of 2% or More Among Aripiprazole-Treated Patients and Greater than Placebo in Short-Term, Placebo-Controlled Trials

Table 2 enumerates the pooled incidence, rounded to the nearest percent, of treatment-emergent adverse events that occurred during acute therapy (up to 6 weeks in schizophrenia and up to 3 weeks in bipolar mania), including only those events that occurred in 2% or more of patients treated with aripiprazole (doses \(\times \) 2mg/day) and for which the incidence in patients treated with aripiprazole was greater than the incidence in patients treated with placebo in the combined dataset.

Table 2: Treatment-Emergent Adverse Events in Short-Term, Placebo-Controlled Trials

	Percentage of Patien	Percentage of Patients Reporting Eventa	
Body System Adverse Event	Aripiprazole (n=1523)	Placebo (n=849)	
Body as a Whole		•	
Headache	31	26	
Asthenia	8	7	
Accidental Injury	5	4	
Peripheral Edema	2	1	
Cardiovascular System			
Hypertension	2	1	
Digestive System			
Nausea	16	12	
Dyspensia	15	13	
Vomiting	11	6 7	
Constipation	11	7	
Musculoskeletal System			
Myalgia	4	3	
Nervous System			
Agitation	25	24	
Anxiety	20	17	
Insomnia	20	15	
Somnolence	12	8	
Akathisia	12	5	
Lightheadedness	11	8	
Extrapyramidal Syndrome	6	4	
Tremor	4	3	
Increased Salivation	3	1	
Respiratory System			
Pharyngitis	4	3 3 2	
Rhinitis	4	3	
Coughing	3	2	
Special Senses			
Blurred Vision	3	1	

^a Events reported by at least 2% of patients treated with aripiprazole, except the following events, which had an incidence equal to or less than placebo: abdominal pain, back pain, dental pain, diarrhea, dry mouth, anorexia, psychosis, hypertonia, upper respiratory tract infection, rash, vaginitis', dysmenorrhea.

Percentage based on gender total.

An examination of population subgroups did not reveal any clear evidence of differential adverse event incidence on the basis of age, gender, or race.

Dose-Related Adverse Events

Dose response relationships for the incidence of treatment-emergent adverse events were evaluated from four trials in patients with schizophrenia comparing various fixed doses (2, 10, 15, 20, and 30 mg/day) of aripiprazole to placebo. This analysis, stratified by study, indicated that the only adverse event to have a possible dose response relationship, and then most prominent only with 30 mg, was somnolence (placebo, 7.7%; 15 mg, 8.7%; 20 mg, 7.5%; 30 mg, 15.3%).

7.7%; 15 mg, 8.7%; 20 mg, 7.5%; 30 mg, 15.3%).

Extrapyramidal Symptoms
In the short-term, placebo-controlled trials of schizophrenia, the incidence of reported EPS for aripiprazole-treated patients was 6% vs. 6% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of reported EPS-related events excluding events related to akathisia for aripiprazole-treated patients was 17% vs. 12% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of akathisia-related events for aripiprazole-treated patients was 15% vs. 4% for placebo. Objectively collected data from those trials was collected on the Simpson Angus Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia) and the Assessments of Involuntary Movement Scales (for dyskinesias). In the schizophrenia trials, the objectively collected data did not show a difference between aripiprazole and placebo, with the exception of the Barnes Akathisia Scale (aripiprazole, 0.05; placebo, -0.05). In the bipolar mania trials, the Simpson Angus Rating Scale and the Barnes Akathisia Scale showed a significant difference between aripiprazole and placebo (aripiprazole, 0.61; placebo, -0.03 and aripiprazole, 0.25; placebo, -0.06). Changes in the Assessments of Involuntary Movement Scales were similar for the aripiprazole and placebo groups.

groups.

Similarly, in a long-term (26-week), placebo-controlled trial of schizophrenia, objectively collected data on the Simpson Angus Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia), and the Assessments of Involuntary Movement Scales (for dyskinesias) did not show a difference between aripiprazole and placebo.

Laboratory Test Abnormalities

A between group comparison for 3- to 6-week, placebo-controlled trials revealed no medically important differences between the aripiprazole and placebo groups in the proportions of patients experiencing potentially clinically significant changes in routine serum chemistry, hematology, or urinalysis parameters. Similarly, there were no aripiprazole/placebo differences in the incidence of discontinuations for changes in serum chemistry, hematology, or urinalysis.

nentational or drindysts.

In a long-term (26-week), placebo-controlled trial there were no medically important differences between the aripiprazole and placebo patients in the mean change from baseline in prolactin, fasting glucose, triglyceride, HDL, LDL, and total cholesterol measurements.

Weight Gain

In 4- to 6-week trials in schizophrenia, there was a slight difference in mean weight gain between aripiprazole and placebo patients (+0.7 kg vs. -0.05 kg, respectively), and also a difference in the proportion of patients meeting a weight gain criterion of ≥7% of body weight [aripiprazole (8%) compared to placebo on patients intesting a weight gain crarefroit of 27% of body weight (arpiprazole (8%) compared to placed (3%)). In 3-week trials in mania, the mean weight gain for aripiprazole and placebo patients was 0.0 kg vs. -0.2 kg, respectively. The proportion of patients meeting a weight gain criterion of ≥7% of body weight was aripiprazole (3%) compared to placebo (2%). Table 3 provides the weight change results from a long-term (26-week), placebo-controlled study of aripiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of ≥7% of body weight relative to baseline, categorized by BMI at baseline:

Table 3: Weight Change Results Categorized by BMI at Baseline: Placebo-Controlled Study in Schizophrenia, Safety Sample

	Bt	VI! <23	BM	1 23-27	BMI	>27
	Placebo	Aripiprazole	Placebo	Aripiprazole	Placebo	Aripiprazole
Mean change from baseline (kg)	-0.5	-0.5	-0.6	-1.3	-1.5	-2.1
% with ≥7% increase BW	3.7%	6.8%	4.2%	5.1%	4.1%	5.7%

Table 4 provides the weight change results from a long-term (52-week) study of aripiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of ≥7% of body weight relative to baseline, categorized by BMI at baseline:

Table 4: Weight Change Results Categorized by BMI at Baseline: Active-Controlled Study in Schizophrenia, Safety Sample

	BMI <23	BMI 23-27	BMI >27
Mean change from baseline (kg)	2.6	1.4	-1.2
% with ≥7% increase BW	30%	19%	8%

ECG Changes

Between group comparisons for a pooled analysis of placebo-controlled trials in patients with schizophrenia or bipolar mania, revealed no significant differences between anipiprazole and placebo in the proportion of patients experiencing potentially important changes in ECG parameters. Aripiprazole was associated with a median increase in heart rate of 5 beats per minute compared to a 1 beat per minute increase among placebo patients.

Additional Findings Observed in Clinical Trials

Additional Findings Observed in Clinical Trials

Adverse Events in Long-Term, Double-Billor, Placebo-Controlled Trials

The adverse events reported in a 26-week, double-blind trial comparing ABiLIFY (aripiprazole) and placebo in patients with schizophrenia were generally consistent with those reported in the short-term, placebo-controlled trials, except for a higher incidence of tremor [9% (13/153) for ABILIFY vs. 19k (2/153) for placebol. In this study, the majority of the cases of tremor were of mild intensity (9/13 mild and 4/13 moderate), occurred early in therapy (9/13 ≤49 days), and were of limited duration (9/13 ≤10 days). Tremor infrequently led to discontinuation (-1%) of ABILIFY in addition, in a long-term (52-week), active-controlled study, the incidence of tremor for ABILIFY was 4% (34/859). A similar adverse event profile was observed in a long-term study in bilonar disorder. study in bipolar disorder.

Other Adverse Events Observed During the Premarketing Evaluation of Aripiprazole
Following is a list of modified COSTART terms that reflect treatment-emergent adverse events as defined in
the introduction to the ADVERSE REACTIONS section reported by patients treated with aripiprazole at
multiple doses ≥2 mg/day during any phase of a trial within the database of 7951 patients. All reported
events are included except those already listed in Table 2, or other parts of the ADVERSE REACTIONS
section, those considered in the WARNINGS or PRECAUTIONS, those event terms which were so general as to
be uniformative quenter record with an incidence of -0.05% and which did not have a publicative be uninformative, events reported with an incidence of \$0.05% and which did not have a substantial probability of being acutely life-threatening, events that are otherwise common as background events, and events considered unlikely to be drug related. It is important to emphasize that, although the events reported occurred

considered unlinkery to be drug related. It is important to enripsace that, authority the during treatment with ariphrazole, they were not necessarily caused by it.

Events are further categorized by body system and listed in order of decreasing frequency according to the following definitions: frequent adverse events are those occurring in at least 1/100 patients (only those not already listed in the tabulated results from placebo-controlled trials appear in this listing); infrequent adverse events are those occurring in 1/100 to 1/1000 patients; rare events are those occurring in fewer than

adverse events are tinuse occurring in 17:00 to 17:00 patients.

Body as a Whole: Frequent – flu syndrome, fever, chest pain, rigidity (including neck and extremity), neck pain, pelvic pain; Infrequent – face edema, suicide attempt, malaise, migraine, chilis, photosensitivity, tightness (including abdomen, back, extremity, head, jaw, neck, and tongue), jaw pain, bloating, enlarged abdomen, chest tightness, throat pain; Rare – moniliasis, head heaviness, throat tightness, Mendelson's

syndrome, heat stroke.

Cardiovascular System: Frequent – tachycardia (including ventricular and supraventricular), hypotension, bradycardia; Infrequent – palpitation, hemorrhage, heart failure, myocardial infarction, cardiac arrest, atrial fibrillation, AV block, prolonged QT interval, extrasystoles, myocardial ischemia, deep vein thrombosis, angina pectoris, palior, cardiopulmonary arrest, philebitis; Aare – bundle branch block, atrial flutter, vasovagal reaction, cardiomegaly, thrombophlebitis, cardiopulmonary failure.

cardinegay, informopriedits, cardiopulmonary railure.

Digestive System: Frequent – nausea and vomiting; Infrequent – increased appetite, dysphagia, gastroenteritis, flatulence, tooth caries, gastritis, gingivitis, gastrointestinal hemorrhage, hemorrhoids, gastroesophageal reflux, periodontal abscess, fecal incontinence, rectal hemorrhage, stomatitis, colitis, tongue edema, cholecystitis, mouth ulcer, oral moniliasis, eructation, fecal impaction, cholelithiasis; Rare

esophagitis, hematemesis, intestinal obstruction, gum hemorrhage, hepatitis, peptic ulcer, glossitis, melena,

duodenal ulcer, cheilitis, hepatomegaly, pancreatitis.

Endocrine System: Infrequent – hypothyroidism; Rare – goiter, hyperthyroidism.

Hemic/Lymphatic System: Frequent – ecchymosis, anemía; Infrequent – hypochromic anemía, leukocytosis, leukopenía (including neutropenía), lymphadenopathy, eosinophilia, macrocytic anemía; Rare –

thrombocythemia, thrombocytopenia, petechiae.

Metabolic and Nutritional Disorders: Frequent – weight loss, creatine phosphokinase increased, dehydration; Infrequent – edema, hyperglycemia, hypercholesteremia, hypokalemia, diabetes mellitus, hypoglycemia, hyperlipemia, SGPT increased, thirst, BUN increased, hyponatremia, SGOT increased, creatinine increased, cyanosis, alkaline phosphatase increased, bilirubinemia, iron deficiency anemia hyperkalemia, hyperuricemia, obesity; Rare - lactic dehydrogenase increased, hypernatremia, gout, hypoglycemic reaction.

Musculoskeletal System: Frequent – muscle cramp; Infrequent – arthralgia, myasthenia, arthrosis, bone pain, arthritis, muscle weakness, spasm, bursitis, myopathy; Rare – rheumatoid arthritis, rhabdomyolysis, tendonitis, tenosynovitis.

Nervous System: Frequent - depression, nervousness, schizophrenic reaction, hallucination, hostility, confusion, paranoid reaction, suicidal thought, abnormal gait, manic reaction, delusions, abnormal dream; Infrequent – emotional lability, twitch, cogwheel rigidity, impaired concentration, dystonia, vasodilation, paresthesia, impotence, extremity tremor, hypesthesia, vertigo, stupor, bradykinesia, apathy, panic attack, decreased libido, hypersomnia, dyskinesia, manic depressive reaction, ataxia, visual hallucination, cerebrovascular accident, hypokinesia, depersonalization, impaired memory, delirium, dysarthria, tardive dyskinesia, amnesia, hyperactivity, increased libido, myocionus, restless leg, neuropathy, dysphoria, hyperkinesia, cerebral ischemia, increased reflexes, akinesia, decreased consciousness, hyperesthesia, slowed thinking; Rare - blunted affect, euphoria, incoordination, oculogyric crisis, obsessive thought, hypotonia,

buccoglossal syndrome, decreased reflexes, derealization, intracranial hemorrhage.

**Respiratory System: Frequent - sinusitis, dyspnea, pneumonia, asthma; Infrequent - epistaxis, hiccup, laryngitis, aspiration pneumonia; *Rare* – pulmonary edema, increased sputum, pulmonary embolism, hypoxia, respiratory failure, apnea, dry nasal passages, hemoptysis.

Skin and Appendages: Frequent - skin ulcer, sweating, dry skin; Infrequent - pruritus, vesiculobullous rash, acne, eczema, skin discoloration, alopecia, seborrhea, psoriasis; Rare - maculopapular rash, exfoliative

Special Senses: Frequent – conjunctivitis; Infrequent – ear pain, dry eye, eye pain, tinnitus, cataract, otitis media, altered taste, biepharitis, eye hemorrhage, deafness; Rare – diplopia, frequent blinking, ptosis, otitis

externa, amblyopia, photophobia.

Urogenital System: Frequent – urinary incontinence; Infrequent – urinary frequency, leukorrhea, urinary retention, cystitis, hematuria, dysuria, amenorrhea, vaginal hemorrhage, abnormal ejaculation, kidney failure, vaginal monillasis, urinary urgency, gynecomastia, kidney calculus, albuminuria, breast pain, urinary burning; Rare – nocturia, polyuria, menorrhagia, anorgasmy, glycosuria, cervicitis, uterus hemorrhage, female lactation, urolithiasis, priapism.

Other Events Observed During the Postmarketing Evaluation of Aripiprazole
Voluntary reports of adverse events in patients taking aripiprazole that have been received since market introduction and not listed above that may have no causal relationship with the drug include rare occurrences of allergic reaction (e.g., anaphylactic reaction, angioedema, laryngospasm, pruritus, or urticaria).

DRUG ARUSE AND DEPENDENCE

Controlled Substance

ABILIFY (aripiprazole) is not a controlled substance.

Abuse and Dependence

Aripiprazole has not been systematically studied in humans for its potential for abuse, tolerance, or physical dependence. In physical dependence studies in monkeys, withdrawal symptoms were observed upon abrupt cessation of dosing. While the clinical trials did not reveal any tendency for any drug-seeking behavior, these observations were not systematic and it is not possible to predict on the basis of this limited experience the extent to which a CNS-active drug will be misused, diverted, and/or abused once marketed. Consequently, patients should be evaluated carefully for a history of drug abuse, and such patients should be observed closely for signs of ABILIFY misuse or abuse (e.g., development of tolerance, increases in dose, drug-seeking

OVERDOSAGE

MedDBA terminology has been used to classify the adverse events.

Human Experience

A total of 76 cases of deliberate or accidental overdosage with aripiprazole have been reported worldwide. These include overdoses with aripiprazole alone and in combination with other substances. No fatality was reported from these cases. Of the 44 cases with known outcome, 33 recovered without sequelae and one recovered with sequelae (mydriasis and feeling abnormal). The largest known acute ingestion with a known outcome involved 1080 mg of aripiprazole (36 times the maximum recommended daily dose) in a patient who fully recovered. Included in the 76 cases are 10 cases of deliberate or accidental overdosage in children (age 12 and younger) involving aripiprazole ingestions up to 195 mg with no fatalities.

Common adverse events (reported in at least 5% of all overdose cases) reported with aripiprazole overdosage (alone or in combination with other substances) include vomiting, somnolence, and tremor. Other clinically important signs and symptoms observed in one or more patients with aripiprazole overdoses (alone or with other substances) include acidosis, aggression, aspartate aminotransferase increased, atrial fibrillation, bradycardia, coma, confusional state, convulsion, blood creatine phosphokinase increased, depressed level of consciousness, hypertension, hypokalemia, hypotension, lethargy, loss of consciousness, QRS complex prolonged, QT prolonged, pneumonia aspiration, respiratory arrest, status epilepticus, and tachycardia

Management of Overdosage
No specific information is available on the treatment of overdose with aripiprazole. An electrocardiogram should be obtained in case of overdosage and, if QTc interval prolongation is present, cardiac monitoring should be instituted. Otherwise, management of overdose should concentrate on supportive therapy, maintaining an adequate airway, oxygenation and ventilation, and management of symptoms. Close medical supervision and monitoring should continue until the patient recovers.

Charcoal: In the event of an overdose of ABILIFY, an early charcoal administration may be useful in partially preventing the absorption of aripiprazole. Administration of 50 g of activated charcoal, one hour after a single 15-mg oral dose of aripiprazole, decreased the mean AUC and C_{max} of aripiprazole by 50%. Hemodialysis: Although there is no information on the effect of hemodialysis in treating an overdose with aripiprazole, hemodialysis is unlikely to be useful in overdose management since aripiprazole is highly bound

Tablets manufactured by Otsuka Pharmaceutical Co., Ltd., Tokyo, 101-8535 Japan or Bristol-Myers Squibb Company, Princeton, NJ 08543 USA
Orally disintegrating tablets manufactured by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA Oral Solution manufactured by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA Distributed and marketed by Otsuka America Pharmaceutical, Inc., Rockville, MD 20850 USA Marketed by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA US Patent Nos: 5,006,528 and 6,977,257



Otsuka America Pharmaceutical, Inc. Rockville, MD 20850 U.S.A.

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ARICEPT helps patients be more like themselves longer™

- Helped keep patients in the community for more than 5 years¹*†
- Is proven effective in cognition, function, and behavior²⁻⁵
- Caregivers spend less time assisting patients with everyday activities⁶
- Established safety and tolerability
- * Results from an observational follow-up of nursing home placement in mild to moderate AD patients (MMSE 10-26) previously enrolled in 1 of 3 randomized, double-blind, placebo-controlled trials with open-label extension phases.
- [†] As with all studies of this type, results may be attributable to various factors. ARICEPT treatment was one such factor.

ARICEPT is indicated for mild to moderate dementia of the Alzheimer's type.

The most common adverse events in clinical trials with ARICEPT were nausea, diarrhea, insomnia, vomiting, muscle cramps, fatigue, and anorexia. In clinical trials, syncopal episodes have been reported (2% for ARICEPT versus 1% for placebo). Cholinesterase inhibitors have the potential to increase gastric acid secretion. Patients at risk for developing ulcers, including those receiving concurrent NSAIDs, should be monitored closely for gastrointestinal bleeding.

Clinical studies of ARICEPT have shown no increase, relative to placebo, in the incidence of either peptic ulcer disease or gastrointestinal bleeding.

Please see brief summary of prescribing information on adjacent page.





References: 1. Geldmacher DS, Provenzano G, McRae T, Mastey V, Ieni JR. Donepezil is associated with delayed nursing home placement in patients with Alzheimer's disease. *J Am Geriatr Soc.* 2003;51: 937-944. 2. Winblad B, Engedal K, Soininen H, et al., and the Donepezil Nordic Study Group. A 1-year, randomized, placebo-controlled study of donepezil in patients with mild to moderate AD. *Neurology.* 2001;57: 481-483. 4. Rogers SL, Doody RS, Moris JC, et al., for the "312" Study Group. A 1-year, placebo-controlled preservation of function survival study of donepezil in AD patients. *Neurology.* 2001;57:481-483. 4. Rogers SL, Doody RS, Mohs RC, Friedhoff LT, and the Donepezil Study Group. Donepezil Study Group. An alzheimer sleesaes: a 15-week, double-blind, placebo-controlled study. *Arch Intern Med.* 1998;158:1021-1031. 5. Rogers SL, Farlow MR, Doody RS, Mohs R, Friedhoff LT, and the Donepezil Study Group. A 24-week, double-blind, placebo-controlled study. *Arch Intern Med.* 1998;158:1021-1031. 5. Rogers SL, Farlow MR, Doody RS, Mohs R, Friedhoff LT, and the Donepezil Study Group. A 24-week, double-blind, placebo-controlled study is the properation of the propezil MSD Study Investigators Group. Efficacy of donepezil on maintenance of activities of daily living in patients with moderate to severe Alzheimer's disease and the effect on caregiver burden. *J Am Geriatr* Soc. 2003;51:737-744.

ARICEPT® ODT (Donepezil Hydrochloride) Orally Disintegrating Tablets
Brief Summary—see package insert for full prescribing information. INDICATIONS AND USAGE ARICEPT® is indicated for the treatment
of mild to moderate dementia of the Alzheimer's type. CONTRAINDICATIONS ARICEPT® is contraindicated in patients with known hypersensitivity to donepezil hydrochloride or to piperidine derivatives. WARNINGS Anesthesia: ARICEPT rypersensitivity to outpear injuriorities of piperiorities derives. Wahnings Ariestnessa: Arice-Fry & a cholimisestinibilitor, is likely to exaggerate succirythohior-bye muscle relaxation during anesthesia: Cartificavascular Conditions: Because of their pharmacological action, cholinesterase inhibitors may have vagotonic effects on the sinoatrial and atrioventricular nodes. This effect may manifest as bradycardia or heart block in patients both with and without known underlying cardiac conduction abnormalities. Syncopa episodes have been reported in association with the use of ARICEPT®. **Gastrointestinal Conditions:** Through their primary action, cholinesterase inhibitors may be expected to increase gastric acid secretion due to increased cholinergic activity. Therefore, patients should be monitored closely for symptoms of active or occult gastrointestinal bleeding, especially those at increased risk for developing ulcers, e.g., those with a history of ulcer disease or those receiving concurrent nonsteroidal anti-inflammatory drugs (NSAIDS). Clinical studies e.g., unserwin a risusy or uncer usases or uncer ecerying concerned interested an air-main manuary urige (ixsaucs). Crimical suitage of ARICEPT® have shown no increase, relative to plazebo, in the incidence of either peptic ulcer disease or gestrointestinal blerding. ARICEPT®, as a predictable consequence of its pharmacological properties, has been shown to produce diarrhea, nausea and vomiting. These effects, when they occur, appear more frequently with the 10 mg/day dose than with the 5 mg/day dose. In most cases, these effects have been mild and transient, sometimes lasting one to three weeks, and have resolved during continued use of ARICEPT®. Genitumiany: Although not observed in clinical trials of ARICEPT®, cholinomimetics may cause bladder outflow obstruction. Neurological Conditions: Seizures: not observed in clinical trials of ARICEPT®, cholinomiments may cause pander outflow obstruction. Neutrological Conditions: Searce Cholinomiments are believed to have some potential to cause generalized convulsions. However, seazure activities may be a manifestation of Alzheimer's Disease. Pulmonary Conditions: Because of their cholinomimetic actions, cholinesterase inhibitors should be prescribed with care to patients with a history of asthma or obstructive pulmonary disease. PRECAUTIONS Drug-Drug Interactions (see Clinical Pharmacology. Clinical Pharmacolinetics: Drug-drug Interactions) Effect of ARICEPT® on the Metabolism of Other Drugs: No in vivo clinical trials have investigated the effect of ARICEPT® on the clearance of drugs metabolised by CYP 2D6 (e.g. imipramine). However, in vitro studies show a low rate of binding to these enzymes (manifestations) and the proposal of the control of K. about 50-130 uM), that, given the therapeutic plasma concentrations of donepezil (164 nM), indicates little likelihood of interference Whether ARICEPT® has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential of ARICEPT® for interaction with theophylline, cimetidine, warfarin, digoxin and keloconazole. No effects of ARICEPT® on the pharmacokinetics of these drugs were observed. Effect of Other Drugs on the Metabolism of ARICEPT®: Keloconazole and quinidine, inhibitors of CYP450, 3A4 and 2D6, respectively, inhibit donepeal metabolism in vitro. Whether there is a clinical effect of quinidine is not known. In a 7-day crossover study in 18 healthy volunteers, ketoconazole (200 mg q.d.) increased mean donepezil (5 mg q.d.) concentrations (AUC_{0.24} and C_{max}) by 36%. The clinical relevance of this increase in concentration is unknown. Inducers of CYP 2D6 and CYP 3A4 (e.g., phenytoin carbamazepine, dexamethasone, rifampin, and phenobarbital) could increase the rate of elimination of ARICEPT® Formal pharmacokinetic studies demonstrated that the metabolism of ARICEPT® is not significantly affected by concurrent administration of digoxin or cimetidine. Use with Anticholinergias: Seause of their mechanism of action, cholinesterase inhibitors have the potential to intelled with the activity of anticholinergia: Seause of their mechanism of action, cholinesterase inhibitors have the potential to intelled with the activity of anticholinergic medications. Use with Cholinomimetics and Other Cholinesterase Inhibitors: A synergistic effect may be expected when cholinesterase inhibitors are given concurrently with succinylcholine, similar neuromuscular blocking agents or cholinergic agonists when cholmestease inhibitors are given concurrently with succeptable productions are given concurrently approached to be such as belfancion. Carcinogenesis, Mutagenesis, Impairment of Fertility No evidence of a carcinogenic potential was obtained in an 88-week carcinogenicity study of donepezil hydrochloride conducted in CD-1 mice at doses up to 180 mg/kg/day (approximately 90 times the maximum recommended human dose on a mg/m² basis), or in a 104-week carcinogenicity study in Sprague-Dawley rats at doses up to 30 mg/kg/day (approximately 30 times the maximum recommended human dose on a mg/m² basis). Donepezil was not mutagenic in the Arnes reverse mutation assay in bacteria, or in a mouse lymphoma forward mutation assay in vitro. In the chromosome abernation test in cultures of Chinese hamster lung (CHL) cells, some clastogenic effects were observed. Donepezil was not clastogenic in the *in* vivormouse micronucleus test and was not clastogenic releasing were userved. Disciplinations was not clastogenic in the *in* vivormouse micronucleus test and was not genotoxic in an *in* vivorunscheduled DNA synthesis assay in rats. Donepezil had no effect on fertility in rats at doses up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m² basis), **Pregnancy Pregnancy Category C**: Teratology studies conducted in pregnant rats at doses up to 16 mg/kg/day (approximately 13 times the maximum recommended human dose on a mg/m² basis) and in pregnant rabits at doses up to 10 mg/kg/day (approximately 16 times the maximum recommended human dose on a mg/m² basis) did not disclose any evidence for a teratogenic potential (approximately to dims be inhabitual in inhomination and an observal angle bash) guild be associated and established by of donepezil. However, in a study in which pregnant rats were given up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m² basis) from day 17 of gestation through day 20 postpartum, there was a slight increase in still births and a slight decrease in pup survival through day 4 postpartum at this dose; the next lower dose tested was 3 mg/kg/day. There are no adequate or well-controlled studies in pregnant women. ARICEPT® should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Nursing Mothers it is not known whether donepezil is excreted in human breast milk. ARICEPT® has no indication for use in nursing mothers. Pediatric Use There are no adequate and well-controlled trials to document the safety and efficacy of ARICEPT® use in nursing mothers. Pediatric Use There are no adequate and well-controlled trials to document the sately and efficacy of ARICEPT in any illness occurring in children. Gerlatric Use Alzheimer's disease is a discorder cocurring printingly in Individuals over 55 years of age. The mean age of the patients enrolled in the clinical studies with ARICEPT® was 73 years; 80% of these patients were between 65 and 84 years old and 49% of the patients were at or above the age of 75. The efficacy and safety data presented in the clinical trials section were obtained from these patients. There were no clinically significant differences in most adverse events reported by patient groups ≥65 years old and <65 years old. ADVERS REACTIONS Adverse Events Leading to Discontinuation The rates of discontinuation from controlled clinical trials of ARICEPT® due to adverse events for the ARICEPT® 5 myddy treatment groups were compatible to those of placebo-treatment groups at approximately 5%. The rate of discontinuation of patients who received 7-day escalations from 5 myddy. to 10 mg/day, was higher at 13%. The most common adverse events leading to discontinuation, defined as those occurring in at least 2% of patients and at twice the incidence seen in placebo patients, are shown in Table 1.

Table 1. Most Frequent Adverse Events Leading to Withdrawal

noni controlled clinical mais by bose group			
Dose Group	Placebo	5 mg/day ARICEPT®	10 mg/day ARICEPT®
Patients Randomized Event/% Discontinuing	355	350	315
Nausea	1%	1%	3%
Diarrhea	0%	<1%	3%
Vomiting	.10/	.10/	20/.

Most Frequent Adverse Clinical Events Seen in Association with the Use of ARICEPT® The most common adverse events, defined as those occurring at a frequency of at least 5% in patients receiving 10 mg/day and twice the placebo rate, are largely precided by ARICEPT® 5. continorminate feeters. These include nausse, diamenta, insomina; womiting, muscle carmy, fatigue and anorexia. These adverse events were often of mild intensity and transient, resolving during continued ARICEPT® treatment without the need for dose modification. There is evidence to suggest that the frequency of these common adverse events may be affected by the rate of tritration. There is evidence to suggest that the frequency of these common adverse events may be affected by the rate of tritration adverse events with 269 patients who received placebo in the 15 and 30-week studies. These patients were titrated to a dose of 10 mg/day over a 6-week period. The rates of common adverse events were lower than those seen in patients tritrated to 10 mg/day over one week in the controlled clinical trials and were comparable to those seen in patients on 5 mg/day. See Table 2 for a comparison of the most common adverse events following one and six week titration regimens.

Table 2. Comparison of Rates of Adverse Events in Patients Titrated to 10 mg/day Over 1 and 6 Weeks

	No tit	ration	One week titration	Six week titration
Adverse Event	Placebo (n=315)	5 mg/day (n=311)	10 mg/day (n=315)	10 mg/day (n=269)
Nausea	6%	5%	19%	6%
Diarrhea	5%	8%	15%	9%
Insomnia	6%	6%	14%	6%
Fatigue	3%	4%	8%	3%
Vomiting	3%	3%	8%	5%
Muscle cramps	2%	6%	8%	3%
Anorexia	2%	3%	7%	3%

Adverse Events Reported in Controlled Trials The events cited reflect experience gained under closely monitored conditions of clinical trials in a highly selected patient population. In actual clinical practice or in other clinical trials, these frequency estimates may not applicate be conditioned to use, reporting behavior, and the kinds of patients treated may differ. Table 3 lists treatment emergent signs and symptoms that were reported in at least 2% of patients in placebo-controlled trials who received ARICEPT® and for which the rate of occurrence was greater for ARICEPT® assigned than placebo assigned patients. In general, adverse events occurred more frequently in female patients and with advencing age.

Table 3. Adverse Events Reported in Controlled Clinical Trials in at Least 2% of Patients Receiving ARICEPT® and at a Higher Frequency than Placebo-treated Patients

Body System/Adverse Event	Placebo (n=355)	ARICEPT® (n=747)
Percent of Patients with any Adverse Event	72	74
Body as a Whole		
Headache	9	10
Pain, various locations	8	9
Accident	6	7
Fatigue	3	5
Cardiovascular System		
Syncope	1	2
Digestive System		
Nausea	6	11
Diarrhea	5 3 2	10
Vomiting	3	5
Anorexia	2	4
Hemic and Lymphatic System		
Ecchymosis	3	4
Metabolic and Nutritional Systems		
Weight Decrease	1	3
Musculoskeletal System		
Muscle Cramps	2	6
Arthritis	1	2
Nervous System		
Insomnia	6	9
Dizziness	6	8
Depression	<1	3
Abnormal Dreams	0	8 3 3 2
Somnolence	<1	2
Urogenital System		
Frequent Urination	1	2
Other Adverse Events Observed During Clinical Trials	ARICEPT® has been admir	istored to over 1700 i

Other Adverse Events Observed During Clinical Trials ARICEPT® has been administered to over 1700 individuals during clinical trials worldwide. Approximately 1200 of these patients have been treated for at least 3 months and more than 1000 patients have been treated for at least 6 months. Controlled and uncontrolled trials in the United States included approximately 900 patients. In regards to the highest dose of 10 mg/day, this population includes 650 patients treated for 3 months, 475 patients treated for 6 months and 116 patients treated for over 1 year. The range of patient exposure is from 1 to 1214 days. Treatment emergent signs and symptoms that occurred during 3 controlled clinical trials and two open-label trials in the United States were recorded as adverse events by the clinical investigators using terminology of their own choosing. To provide an overall estimate of the proportion of individuals having similar types of events, the events were grouped into a smaller number of standardized categories using a modified COSTART dictionary and event frequencies were calculated across all studies. These categories are used in the listing below. The frequencies represent the proportion of 900 patients from these trials who experienced that event while receiving ARICEPT® All adverse events occurring at least twice are included, except for those already listed in Tables 2 or 3, COSTART terms to general to be informative, or events esculed a relative and entitled a charged listed in Tables 2 or 3, COSTART terms to general to be informative, or events less likely to be drug caused. Events are classified by body system and listed using the following definitions: frequent adverse events—those occurring in at least 1/100 patients; infrequent adverse events—those occurring in at least 1/100 patients, infrequent and in most cases were observed at a similar frequency in placebo-treated patients in the controlled studies. No important additional adverse and infloot cases were used as disminial inequency in place-conceasing patients in the control extousies not important account and events were seen in studies conducted outside the United States. **Body as a Whole:** Frequent: influenza, chest pain, toothache, Infrequent: fever, edema face, periorbital edema, hernia hiatal, abscess, cellulitis, chills, generalized coldness, head fullness, listlessness. **Cardiovascular System:** Frequent: Phypertension, vascodialition, atrial fibrillation, hot flashes, hypotension, in the quent: anging pectoris, postural hypotension, mycoardial infraction, AV block (first degree), congestive heart fallure, arteritis, bradycardia, peripheral pacular disease, supraventricular tachycardia, deep vein thrombosis. **Digestive System:** Frequent: fecal incontinence, gastrointestinal bleeding, bloating, epigastric pain; Infrequent: eructation, gingivitis, increased appetile, flatulence, periodontal abscess, cholelithiasis, diverticulitis, drooling dry mouth, lever sore, gastritis, irritable colon, tongue edema, epigastric distress, gastroenteritis, increased transaminases, hemorrhoids ileus, increased thirst, jaundice, melena, polydipsia, duodenal ulcer, stomach ulcer. Endocrine System: Infrequent: diabetes mellitus, goiter. Hemic and Lymphatic System: Infrequent: anemia, thrombocythemia, thrombocytopenia, eosinophilia, erythrocytopenia. Metabolic and Nutritional Disorders: Frequent: dehydration; Infrequent: gout, hypokalemia, increased creatine kinase, hyperglycemia, weight increase, increased lactate dehydrogenase. Musculoskeletal System: Frequent: bone fracture; Infrequent: muscle weakness, muscle fasciculation. Nervous System: Frequent: delusions, tremor, irritability, paresthesia, aggression, vertigo, ataxia, increased libido restlessness, abnormal crying, nervousness, aphasia; Infrequent: cerebrovascular accident, intracranial hemorrhage, transient ischemic altack, emotional lability, neuralgia, coldness (localized), musole spasm, dysphoria, gait abnormality, hypertonia, hypokinasia, neurodemaltis, numbness (localized), paranoia, dysarthria, dysphasia, hostility, decreased libido, melancholia, emotional withdrawal, nystagmus, pacing.

Respiratory System: Frequent: dyspnea, sore throat, bronchitis; Infrequent: epistaxis, post nasal drip, pneumonia, hyperventilation, pulmonary congestion, wheezing, hypoxia, pharyngitis, pleurisy, pulmonary collapse, sleep apnea, snoring. **Skin and Appendages**Frequent: pruritus, diaphoresis, urticaria; Infrequent: dermalitis, erythema, skin discoloration, hyperkeratosis, alopecia, fungal dermalitis Frequent pruritus, diaphoresis, urticaria, infraquent dermatitis, erythema, skin discoloration, hyperkeratosis, alopecia, fungal dermatitis, erperse zoster, hirsuitsm, skin striae, night sweats, skin ulcer. Special Senses: Frequent: cataract, eye irritation, vision blurred, infraquent: dry eyes, glaucoma, earache, tinnitus, blephantis, decreased hearing, retinal hemorrhage, citilis externa, otitis media, bad tasle, conjunctival hemorrhage, ear buzzing, motion sickness, spots before eyes. Urogenital System: Frequent: urinary incontinence, nocturia. Infraquent: dysuria, hematuria, urinary urgenor, metrorrhagia, cystitis, enuresis, prostate hypertrophy, pyelonephritis, inability to empty bladder, breast fibroadenosis, fibrocystic breast, mastitis, pyruria, renal failure, vagninis. Postintroduction Reports Voluntary reports of adverse events temporally associated with ARICEPT® that have been received since market introduction that are not listed above, and that there is inadequate data to determine the causal relationship with the drug include the following: abdominal pain, agitation, cholecystitis, confusion, repulsions by the purcipation sequence purcipation are purcipation. convulsions, hallucinations, heart block (all types), hemolytic anemia, hepatitis, hyponatremia, neuroleptic malignant syndrome, pancrealitis, and rash. **OVERDOSAGE Because strategies for the management of overdose are continually evolving, it is** advisable to contact a Poison Control Center to determine the latest recommendations for the management of an overdose of any drug. As in any case of overdose, general supportive measures should be utilized. Overdosage with cholinesterase inhibitors can result in cholinergic crisis characterized by severe nausea, vorniting, salivation, sweating, bradycardia, hypotension, respiratory depression, collapse and convulsions. Increasing muscle weakness is a possibility and may result in death if respiratory muscles are involved. Tertiary anticholinergics such as atropine may be used as an antidote for ARICEPT® overdosage. Intravenous atropine sulfate titrated to effect is recommended: an initial dose of 1.0 to 2.0 mg (V with subsequent doses based upon clinical response. Alypical responses in blood pressure and heart rate have been reported with other cholinomimetics when co-administered with quaternary anticholinergics such as glycopyrrolate and rear late level using profession under Commission when Covariant seed with quadrate by anticulturing social suppropriets with control that it is not known whether ARICEPT® and/or its metabolities can be removed by dialysis (hemodialysis, perinoreal dialysis, or hemofiltration). Dose-related signs of toxicity in animals included reduced spontaneous movement, prone position, staggering gait, lacrimation, cloric convulsions, depressed respiration, salivation, microis, temorrs, fasciculation and lower body surface temperature. **DOSAGE AND ADMINISTRATION** The dosages of ARICEPT® shown to be effective in controlled clinical trials are 5 mg and 10 mg administered once per day. The higher dose of 10 mg did not provide a statistically significantly greater clinical benefit than 5 mg. There is a suggestion, however based upon order of group mean scores and dose trend analyses of data from these clinical trials, that a daily dose of 10 mg of ARICEPT[©] based upon foued of upon finear source and udoes return alrayes or load in or misce criminar institute, and a daily ouse of fining or whole; in might provide additional benefit for some patients. Accordingly, whether or not to employ a dose of 10 mg is a matter of prescriber and patient preference. Evidence from the controlled trials indicates that the 10 mg dose, with a one week titration, is likely to be associated with a higher incidence of cholinergic adverse events than the 5 mg dose. In open label trials using a 6 week titration, the frequency of these same adverse events was similar between the 5 mg and 10 mg dose groups. Therefore, because steady state is not achieved for 15 days and because the incidence of untoward effects may be influenced by the rate of dose escalation, treatment with a dose of 10 mg should not be contemplated until patients have been on a daily dose of 5 mg for 4 to 6 weeks. ARICEPT®/ARICEPT® ODT should be taken in the evening, just prior to retiring. ARICEPT®/ARICEPT® ODT can be taken with or without food. Allow ARICEPT® ODT tablet to dissolve on the tongue and follow with water



Pfizer U.S. Pharmaceutical

200337 Revised February 2005

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Control acute agitation with

GEODON® for Injection | ziprasidone mesylate|

In schizophrenia...

Rapid improvement with low EPS^{1,2}

- Significant control achieved between 15 and 30 minutes* after injection^{1,3}
- Proven advantages over haloperidol IM
 - twice the improvement as measured on the BPRS^{4†}
 - significantly lower incidence of movement disorders^{2‡}
- Smooth transition, with continued improvement, from IM to oral therapy^{2,4}
- May be used concomitantly with benzodiazepines

*In 2 pivotal studies vs control, significance was achieved at 15 minutes (with 10 mg dose) and 30 minutes (with 20 mg dose), respectively.

†In a 7-day, open-label IM-to-oral transition study.

*In a 6-week, open-label IM-to-oral transition study.



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.

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.

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 (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.

LINDICATIONS—GEODON Capsules is indicated for the treatment of schizophrenia and acute manic or mixed episodes associated with bipolar disorder with or without psychotic features. GEODON® (ziprasidone mesylate) for Injection is indicated for acute agitation in achieve benefit and the provided by the p

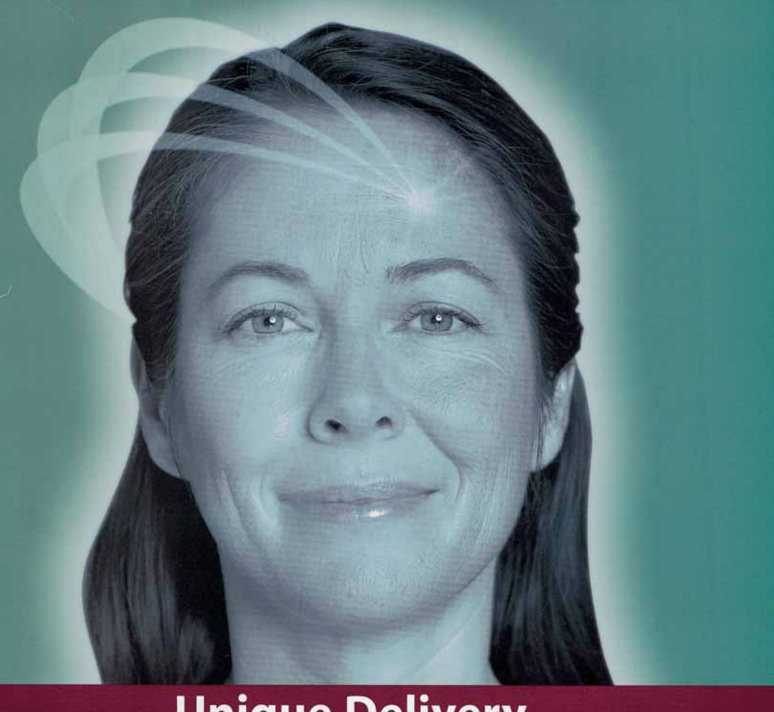
CONTRAINDICATIONS — QT Prolongation: Because of GEODON's dose-related prolongation of the QT interval and the known associate of fatal arrhythmias with 0T prolongation by some other drugs, GEODON is contraindicated in patients with a known history of OT prolongation (including congenital long QT syndrome), with recent acute myocardial infarction, or with uncompensated heart failure (see WARNINGS). Pharmacokinetic/pharmacodynamic studies between GEODON and other drugs that prolong the QT interval have not been performed. An additive effect of GEODON and other drugs that prolong the OT interval cannot be excluded. Therefore, GEODON should not be given with dofetilide, sotalol, quinidine, other Class Ia and III anti-arrhythmics, mesoridazine, thioridazine, chlorpromazine, droperidol, pimozide, sparfloxacin, gatifloxacin, moxifloxacin, halofantrine, mefloquine, pentamidine, arsenic trioxide, levomethadyl acetate, dolasetron mesylate, probucol, or tacrolimus, GEODON is also contraindicated with drugs that have demonstrated QT prolongation as one of their mesylate, productor, accominus, secution is sale contraminated with originating that have employed profit p identification of other drugs that have been consistently observed to prolong the OT, interval. Such drugs should not be prescribed with GEODON. A study directly comparing the OT/OT_c-prolonging effect of GEODON with several other drugs effective in the treatment of eshipotherina was conducted in patient volunteers. The mean increase in OT_c from baseline for GEODON ranged from approximately 9 to 14 mise greater than for four of the comparator drugs (risperidone, olanzapine, quetlapine, and haloperidol), but was approximately 14 misec less than the prolongation observed for third indicate. In this study, the effect of 65000N on 75_length was not augmented by the presence of a metabolic inhibitor (ketoconazole 200 mg bid). In placebo-controlled trials, GEODON increased the OT_interval compared to placebo by approximately 10 misec at the highest recommended daily dose of 160 mg. In clinical trials the electrocardiograms of 2/2988 (0.06%) GEODON patients and 1/440 (0.23%) placebo patients revealed OT_interval trials the potentially clinically relevant threshold of 500 misec. In the GEODON patients, neither case suggested a role of GEODON. Some drugs that prolong the OT/OT_interval have been associated with the occurrence of forsade de pointes and with sudden unexplained death. The relationship of QT prolongation to torsade de pointes is clearest for larger increases (20 mase and greater) but its possible that smaller QT/QT_c prolongations may also increase risk, or increase it in susceptible individuals, such as those with hypokalemia, hypomagnesemia, or genetic predisposition. Although torsade de pointes has not been observed in association with the use of GEODON nypomagnesemia, or genetic previsions no. Almough oursade de polimes has not deem observed in association with the use of LEDUN at recommended doses in premarketing studies, experience is too limited to rule out an increased risk. A study on the property prologing effect of intramuscular GEDDON, with intramuscular haloperidol as a control, was conducted in patient volunteers. In the trial, ECGs were obtained at the time of maximum plasma concentration following buo injections of GEDDON is 50% higher than the recommended therapeutic dose. The mean change in QT₂ from baseline was calculated for each drug using a sample-based correction that removes the effect of heart rate on the QT interval. The mean increase in QT₄ from baseline for GEDDON was 4.6 mese following the first injection and 12.8 msec following the second injection. The mean increase in QT₄ from baseline for haloperidol was 6.0 msec following the second injection. This study, no patient had a QT₄ interval exceeding 500 msec. As with other antipsychotic drugs and placebo, sudden unexplained deaths have been reported in patients taking GEDDON at recommended doses. The premarketing experience for GEDDON did not reveal an excess of mortality for GEDDON compared to other antipsychotic drugs or placebo, but the extent of exposure was limited, especially for the drugs used as active controls and placebo. Nevertheless, GEDDONs larger prolongation of QT₂ length compared to several other antipsychotic drugs raises the possibility that the risk of studden death may be greater for EGEDDON than for other available drugs for treating schizophrenia. The presence of conjects to be considered in deciding among alternative drug products. Certain circumstances may increase the risk of the occurrence of torsade de pointes and/or sudden death in association with the use of drugs that prolong the QT₂ interval, including (1) bradyeardia; (2) hypokalemia or hypomagnessemia; (3) concominatant use of other drugs that prolong the QT₂ interval; and (4) presence of at recommended doses in premarketing studies, experience is too limited to rule out an increased risk. A study evaluating the QT/QT. history of cardiac arrhythmias (see CONTRAINDICATIONS), and see *Drug Interactions* under PRECAUTIONS). It is recommended that patients being considered for GEODON treatment who are at risk for significant electrolyte disturbances, hypokalemia in particular, have baseline serum potassium and magnesium measurements. Hypokalemia (and/or hypomagnesemia) may increase the risk of OT prolongation and arrhythmia. Hypokalemia may result from diuretic therapy, diarrhea, and other causes. Patients with lows erum potassium and/or magnesium should be repleted with those electrolytes before proceeding with treatment. It is essential to periodically monitor serum electrolytes in patients for whom diuretic therapy is introduced during GEODON treatment. Persistently prolonged OT_c intervals may also increase the risk of further prolongation and arrhythmia, but it is not clear that routine screening ECG measures are effective in detecting such patients. Rather, GEODON should be avoided in patients with histories of significant cardiovalar illness, eg, QT prolongation, recent acute myocardial infarction, uncompensated heart failure, or cardiac arrhythmia. GEODON should be discontinued in patients who are found to have persistent OT, measurements >500 msec. *Neurolophic Malignant Syndrome (MMS)*: A hope been pronder in the potentially fatal symptom complexes ymmelines referred thas Neuroleptic Malignant Syndrome (MMS): As been reported in conscious of the properties of the potentially fatal speem reported in conscious of the properties of the propertie potentially fatal symptom complex sometimes referred to as Neuroleptic Malignant Syndrome (NMS) has been reported in association with administration of antipsychotic drugs. The management of MMS should include: (1) immediate discontinuation of antipsychotic drugs and other drugs not essential to concurrent therapy, (2) intensive symptomatic treatment and medical monitoring; and (3) treatment of any concomitant serious medical problems for which specific treatments are available. If a patient requires antipsychotic drug treatment after recovery from NMS, the potential reintroduction of drug therapy should be carefully considered. The patient should be carefully monitored, since recurrences of NMS have been reported. *Tardive Dyskinesia (TD):* A syndrome of potentially irreversible, involuntary, dyskinetic movements may develop in patients undergoing treatment with antipsychotic drugs. Although the prevalence of TD appears to be highest movements may develop in patients undergoing treatment with antipsychotic drugs. Amough the prevaelnce or 10 appears to be ingra-among the elderly, especially elderly women, it is impossible to rely upon prevalence estimates to predict, at the inception of antipsychotic treatment, which patients are likely to develop Tb. If signs and symptoms of Tb appear in a patient on GEODON, drug discontinuations brould be considered. **Pyperglycemia and **Diabetes* **Melitius:**Hyperglycemia-related adverse events, sometimes serious, have 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 abplical antipsychotic should be monitored to symptoms of hyperglycemia. PRECAUTIONS — **General: **Rasti, in premarketing trials, about 5% of GEODON patients developed rash and/or urticaria, with discontinuation of treatment in about one-sixth of these cases. The occurrence of rash was dose related, although the findion moints also be explained the longer exposures in bibber-dose artisents. Swertlo altorsts with rest hat eigens and surrous described. and/or uncaria, with discontinuation of treatment in about one-sixent of these cases. The occurrence or rash was oose related, although finding might also be explained by longer exposure in higher-dose patients. Several patients with rash had signs and symptoms of associated systemic illness, e.g., elevated WBCs. Most patients improved promptly upon treatment with antihistamines or steroids and/or upon discontinuation of ECDODN, and all patients were reported to recover completely. Upon appearance of rash for which an attendate eitoria cannot be identified, GEODON should be discontinued. Orthostatic Hypotension: GEODON may induce orthostatic hypotension associated with disziness, tachycardia, and, in some patients, syncope, especially during the initial dose-liteation period, probably reflecting its c₁-adrenergic antagonist properties. Syncope was reported in 0.5% of GEODON patients. GEODON should be used with particular caution in patients with known cardiovascular disease (history of myocardial infarction or ischemic heart disease, heart failure or conduction abnormalities), cerebrovascular disease or conditions that would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications). Seizures: In clinical trials, seizures occurred in 0.4% of GEODON patients. There were confounding factors that may have contributed to seizures in many of these cases. As with other antipsychotic drugs, GEODON should be used cautiously in patients with a history of seizures or with conditions that potentially lower the seizure threshold, e.g., Atheimer's dementia. Conditions that lower the seizure threshold may be more prevalent in a population of 65 years or older. <u>Dysphagia:</u> Esophageal dysmotility and a spiration have been associated with antipsychotic drug use. Aspiration preumonia is a common cause of morbidity and mortality in elderly patients, in particular those with advanced Atheimer's dementia, and GEDDON and other antipsychotic drugs should be used cautiously in patients at risk for aspiration pneumonia. (See also Boxed WARNING, WARNINGS: Increased Mortality in Elderly Patients with Dementia-Related Psychosis). Hyperprolactinemia: As with other drugs that antagonize dopamine D₂ receptors, GEODON elevates prolactin levels in humans. Tissue culture experiments indicate that approximately one third of human breast cancers are prolactin dependent in vitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancer. Whether clinical studies no repidemiologic studies conducted to date have shown an association between chronic administration of this class of drugs and tumorigenesis in humans; the available evidence is considered too limited to be conclusive at this time. Potential for Cognitive and Motor Impairment. Somnolence was a commonly reported adverse event in ECDOON patients. In the 4- and 6-week placebo-controlled trials, somnolence was reported in 14% of GEODON patients is 1600 patients. Somnolence led to discontinuation in 0.3% of patients in short-term clinical trials. Since GEODON has the potential to impair judgment, thinking, or motor skills, patients should be cautioned about performing activities requiring mental alertness, such as operating a motor vehicle (including automobiles) or operating hazardous machinery until they are reasonably certain that GEODON therapy does not affect them adversely. <u>Praipsim:</u> One case of praipsim was reported in the premarketing database. <u>Body Temperature Regulation</u>: Although not reported with GEODON in premarketing trials, disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents. <u>Suicide</u>: The possibility of a suicide attempt is inherent in psychotic illness and close supervision of high-risk patients should accompany drug hepsay. Because in suicide attempt is inherent in psychotic illness and close supervision of high-risk patients should accompany drug therapy, GEODON prescriptions should be written for the smallest quantity of capsules consistent with good patient management to reduce overdose risk. <u>Use in Patients with Concomitant Illness</u>; Clinical experience with GEODON in patients with certain concomitant systemic illnesses is limited. GEODON has not been evaluated or used to any appreciable extent in patients with a recent history of myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from premarketing clinical studies. Because of the risk of QT_C prolongation and orthostatic hypotension with GEODON, caution should be observed in cardiac patients (see *QT Prolongation and Risk of Sudden Death* in WARNINGS and Orthostatic Hypotension in PRECAUTIONS). Information for Patients: To ensure safe and effective use of GEODON, the

information and instructions in the Patient Information Section should be discussed with patients. Laboratory Tests: Patients being considered for GEODON treatment who are at risk of significant electrolyte disturbances should have baseline serum potassium and magnesium measurements. Low serum potassium and magnesium should be repleted before treatment. Patients who are started on diuretics during GEODON therapy need periodic monitoring of serum potassium and magnesium. Discontinue GEODON in patients who are found to have persistent OT_c measurements >500 msec (see **WARNINGS**). *Drug Interactions*: (1) GEODON should not be used with any drug that prolongs the OT interval. (2) Given the primary CNS effects of GEODON, caution should be used when it is taken in combination with other centrally the U1 merval. (2) Given the primary CNs effects of tecturous, caution should be used when it is taken in comination with other certain artifypacting drugs. (3) Because of its potential for including hypotension, ECDDOM may enhance the effects of certain antihypertansive agents. (4) GEDDOM may antagonize the effects of levodopa and dopamine agonists. <u>Effect of Other Drugs on GEDDOM</u>; Carbamazepine, 200 mg bid for 21 days, resulted in a decrease of dapproximately 35% in the AUC of GEDDOM. **Reboornazele*, a potent inhibitor of CYP3A4, 400 mg dfor 5 days, in creased the AUC and Cm_{max} of GEDDOM by about 35% -40%. **Cimetion, 800 mg of for 2 days, did not affect GEDDOM pharmacokinetics. Population pharmacokinetic analysis of schizophrenic patients in controlled clinical trials has not revealed any clinically significant pharmacokinetic interactions with benztropine, propranolol, or lorazepam. <u>Effect of GEDDOM</u> on <u>Other Drugs</u>; in vitro studies revealed little potential for GEDDOM to therefore with the metabolism of drugs cleared primarily by CYP1A2. CYP2CS, CYP2C19, CYP2D6, and CYP3A4, and little potential for drug interactions with the metabolism of drugs cleared primarily by CYP1A2. CYP2CS, CYP2C19, CYP2D6, and CYP3A4, and little potential for drug interactions with GEODON due to displacement. GEODON 40 mg bid administered concomitantly with ##### 450 mg bid for 7 days did not affect the steady-state level or renal clearance of lithium. GEODON 20 mg bid did not affect the pharmacokinetics of concomitantly administered oral contraceptives ethinyl estradiol (0.03 mg) and evonorgestrel (0.15 mg). Consistent with invitro results, a study in normal healthy volunteers showed that GEODON did not alter the metabolism of dextromethorphan, a CYP2D6 model substrate, to its major metabolite, dextropphan. There was no statistically significant change in the urinary dextromethorphan/dextrophan ratio. Carcinogenesis, Mutagenesis, Impairment of Fertility. Lifetime carcinogenicity studies were conducted with GEODON in Long Evans rats and CD1 mine. In male mice, there was no increase in incidence of tumors relative to controls. In fernale mice there were dose-related increases in the incidences of pituitary gland adenoma and carcinoma, and mammary gland adenocarcinoma at all doses tested. Increases in serum prolactin were observed in a 1-month dietary study in female, but not male, mice. GEODON had no effect on serum prolactin in rats in a 5-week dietary study at the doses that were used in the carcinogenicity study. The relevance for human risk of the findings of prolactin-mediated endocrine tumors in rodents is unknown (see https://https://https:rpcreatable-newfatth-mediated endocrine tumors in rodents is unknown (see https://ht gene mutation assay and the in vitro chromosomal aberration assay in human lymphocytes. <u>Impairment of Fertility</u>. GEODON increased time to copulation in Sprague-Dawley ratis in two fertility and early embryonic development studies at doses of 10 to 160 mg/kg/day (0.5 to 8 times the MRHD of 200 mg/day on a mg/m² basis). Fertility rate was reduced at 160 mg/kg/day (8 times the MRHD on a mg/m² basis). The retility of female rats was reduced. **Pregnancy—** Pregnancy Category C: There are no adequate and well-controlled studies in pregnant women. GEODON should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Labor and Delivery: The effect of GEODON on labor and delivery in humans is unknown. Nursing Mothers: It is not known whether, and if so in what amount, GEODON or its metabolites are excreted in human milk. It is recommended that women receiving GEODON should not breast feed. **Pediatric Use:** The safety and effectiveness of GEODON in pediatric patients have not been established. *Geriatric Use:* Of the approximately 4500 patients treated with GEODON in clinical studies, 2.4% (109) were 65 years of age or over. In general, there was no indication of any different tolerability for GEODON or of reduced clearance of GEODON in the elderly compared to younger adults. Nevertheless, the presence of multiple factors that might increase the pharmacodynamic response to GEODON, or cause poorer tolerance or orthostasis, should lead to consideration of a lower starting dose, slower titration, and careful monitoring during the initial dosing period for some elderly patients. ADVERSE REACTIONS— Adverse Findings Observed in Short-term, Placebo-Controlled Trials: The following findings are based on the short-term placebo-controlled premarketing trials for schizophrenia (a pool of two 6-week, and two 4-week fixed-dose trials) and bipolar mania (a pool of two 3-week flexible-dose trials) in which GEODON was administered in doses ranging from 10 to 200 mg/day. *Adverse Events Associated with Discontinuation:* Schizophrenia: Approximately 4.1% (29/702) of GEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 2.2% (6/273) on placebo. The most common event associated with dropout was rash, including 7 dropouts for rash among GEODON patients (1%) compared to no placebo patients (see **PRECAUTIONS**). Bipolar Mania: Approximately 6.5% (18/279) of GEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 3.7% (5/136) on placebo. The most common events associated with dropout in the GEODON-treated adverse event, complain with about 3 of 18 of 19 in place. The instrument event as sociated with original managed patients were akathisia, anxiety, depression, dizziness, dystonia, rash and vomiting, with 2 dropouts for each of these events among €EDON patients (1%) compared to one placebo patient each for dystonia and rash (1%) and no placebo patients for the remaining adverse events.

Adverse Events at an Incidence ≥ 5% and at Least Twice the Rate of Placebo: The most commonly observed adverse events associated with GEODON in schizophrenia trials were sonnolence (14%) and respiratory tract infection (8%). The most commonly observed adverse events associated with the use of GEODON in bipolar mania trials were somnolence (31%), extrapyramidal symptoms (31%), dizziness (16%), akathisia (10%), abnormal vision (6%), asthenia (6%), and vomiting (5%). The following list enumerates the treatment-emergent (16%), alathisia (10%), ahonomal vision (6%), ashenia (6%), and vomiting (5%). The following list enumerates the treatment-emergent adverse events that occurred unique acute therapy, including only those events that occurred in 2% of GEDDON patients and at greater incidence than in placebo. Schizophrenia: Body as a Whole—asthenia, accidental injury, chest pain. Cardiovascular—tachycardia. Digestive—nausea, constipation, dyspepsia, diarrhea, dry mouth, anorexia. Neurous—extrapyramidal symptoms, somnolenca, aktalika diziness. Respiratory—respiratory tract infection, rhinitis, cough increased. Skin and Appendages—rash, fungal dematitis. Special Senses—abnormal vision. Bipolar Mania: Body as a Whole—headache, asthenia, accidental injury. Cardiovascular—hypertension. Digestive—nausea, diarrhea, dry mouth, vomiting, increased salivation, tongue edema, dysphagia. Musculoskeletal—myolgia. Menson somnolence, extrapyramidal symptoms, dizziness, akathisia, anavely, hypesthesia, speech disorder. Respiratory—pharyngitis, dysponea. Skin and Appendages—fungal dermatitis. Special Senses—abnormal vision. Dase Dependency: An analysis for dose response in the schizophrenia thatis revealed an apparent relation of adverse event to dose for the following: asthenia, postural hypotension, anorexia, dry mouth, increased salivation, arthralgia, anviety, dizziness, dystonia, hypertonia, somnolence, tremor, rhinitis, rash, and abnormal vision. Extrapyramidal Symptoms (EPS): The incidence of reported EPS for GEODON patients in the short-term, placebo-controlled schizophrenia trials was 14% vision. Scale and these Akathisia and scale and the Base Akathisia and Sarka and Base Akathisia and Sarka and the Base Akathisia and Sarka and Base Akat Extrapyraminal Symptoms (EMS): The incidence of reported EMS for EUDUN patients in the short-term, placebo-controlled scrizophrenia trials aws 14% as % for placebo. Objectively collected data from those trials on the Simpson-Angus Raling Scale and the Base Akathisia Scale did not generally show a difference between GEODON and placebo. Vital Sign Changes: GEODON is associated with orthostatic hypotension (see PRECAUTIONS). Weight Cain: In short-term schizophrenia trials, the proportions of patients meeting a weight gain criterion of 27% of body weight were compared, revealing a statistically significantly greater incidence of weight gain for GEODON patients (10%) is placebo patients (4%). A median weight gain for 0.5 kg was observed in GEODON, patients vs 0.0 kg in placebo patients. Weight gain was reported as an adverse event in 0.4% of both GEODON, and placebo patients. During long-term therapy with GEODON, a categoristic field of centures the design of the basic of flow more index (40M) showed the consideration of the content of the centure of the content of the content of the centure of the content of the centure of the centure of the content of the centure of the centu gain was reported as an adverse event in U-4% or blood in EcODOV and placebo placents. Joining long-lent in lendary with Incoord categorization of platents at baseline on the basis of body mass index (EMI) showed the greatest mean weight gain and the highest incidence of clinically significant weight gain and the highest incidence of clinically significant weight gain and the Might platents. There was a mean weight gain of 1.4 kg for patients with a "low" baseline BMI, 0.0 kg for patients with a "normal" BMI, and a 1.3 kg mean weight loss for patients with a "high" BMI. ECG Changes: GEODON is associated with an increase in her OT, interval (see WARNINGS). In schizophrenia trials, GEODON was associated with a mean increase in heart rate of 1.4 beats per minute compared to all beats per minute decrease among placebo patients. Other Adverse Events Observed Unring the Premarketing Evaluation of GEODON: Frequent adverse events are those occurring in at least 1/100 patients; infrequent adverse events are those occurring in 1/100 to 1/1000. patients: rare events are those occurring in fewer than 1/1000 patients. Schizophrenia: <u>Body as a Whole</u>—*Frequent*: abdominal pain, flu syndrome, fever, accidental fall, face edema, chills, photosensitivity reaction, flank pain, hypothermia, motor vehicle accident. <u>Cardiovascular</u> <u>System</u>—*Frequent* tachycardia, hypertension, postural hypotension; *Infrequent*: bradycardia, angina pectoris, atrial fibrillation; *Rare*: first-<u>Ogstein Treguent authyrotation in protein to many proteins in many proteins and a region proteins and a signal proteins and a degree AV block, buileb tranch block, phlebits, pulmonary embolus, cardiomegaly, cerebral infarct, cerebrovascular accident, deep thrombophlebitis, myocardiitis, thrombophlebitis. <u>Digestive System</u>— *Frequent*: anorexia, vomiting; *Infrequent*: rectal hemorrhage, dysphagia, tongue edema; *Ran*e; gum hemorrhage, jaundice, fecal impaction, gamma glutamy transpeptidase increased, hematements.</u> cholestatic jaundice, hepatitis, hepatomegaly, leukoplakia of mouth, fatty liver deposit, melena. Endocrine—Rare: hypothyroidism knieszab Jannuck, prejamus, repartingelay, teuropiana or mount, taty vier deposit, intereista <u>Endocrine</u>—hare Tryboury louiste. Hyperthyroidism, thyroiditis. <u>Hemic and Lymphatic System</u>—*Infrequent*: anemia, ecchymosis, leukocytosis, leukopenia, eosinophilia, lymphadenopathy, *Rare*: thrombocytopenia, hypochromic anemia, lymphocytosis, monocytosis, basophilia, lymphedema, polycythemia, thrombocythemia. <u>Metabolic and Nutritional Disorders</u>—*Infrequent*: thirst, transaminase increased, peripheral edema, hyperglycemia, pympiaueriopaury, Raze untimocoypenia, hypocitromicalenia, gympiocytosis, tasoprojussis, assignia, yrimpieuerina, piotycyterina, thrombooythemia, Metabolic and Nutritional Disorders — Infrequent thirist, transaminase increased, enpiriperal edema, hyperglycemia, creatine phosphokinase increased, alkaline phosphokinase increased, alkaline phosphokinase increased, alkuline phosphokinased, alkuline phosphokinase increased, alkuline phosphokinase in Incidence 19% in Short-Term Fixed-Dose Intramuscular Triols: The following list enumerates the treatment-emergent adverse events that occurred in ≥1% of GEODON patients (in the higher dose groups) and at least twice that of the lowest intramuscular GEODON group. Body as a Whole—headache, injection site pain, asthenia, abdominal pain, flusyndrome, back pain. Cardiovascular—postural hypotension, <u>body as wynue</u>—lieadacke, lijectourise pain, saleinia, abuoriliari pain, systienia, systyntrulie, tacky an tis <u>dallovyascula</u>—pustural pytoelistoin, hypertension, bradycardia, vasodilation. <u>Digestive</u>—nausea, rectal hemorrhage, diarrhea, vomiting, olysepesia, anorexia, constipation, tooth disorder, dry mouth. <u>Nervous</u>—dizziness, arxiety, insomnia, somnolence, akathisia, agitation, extrapyramidal syndrome, hypertonia, cogwheel rigidity, paresthesia, personality disorder, psychosis, speech disorder. <u>Respiratory</u>—rhinitis. <u>Skin and Appendages</u>—frurunculosis, sweating. <u>Urogenital</u>—dysmenorrhea, priapism. **DRUG ABUSE AND DFENDENCE—*Controlled Substance Class*:
GEODON is not a controlled substance. **OVERDOSAGE**—In premarketing trials in over 5400 patients, accidental or intentional overdosage of GEODON was documented in 10 patients. All patients survived without sequelae. In the patient taking the largest confirmed amount (3240 mou), the only complete promoters provided unexample in the patient baking the largest confirmed amount (3240 mou). mg), the only symptoms reported were minimal sedation, slurring of speech, and transitory hypertension (BP 200/75)

References: 1. Daniel DG, Potkin SG, Reeves KR, Swift RH, Harrigan EP. Intramuscular (IM) ziprasidone 20 mg is effective in reducing acute agitation associated with psychosis: a double-blind, randomized trial. Psychopharmacology. 2001;155:128-134. 2. Brook S, Walden J, Benattia I, Siu CO, Romano SJ. Ziprasidone and haloperidol in the treatment of acute exacerbation of schizophrenia and schizoaffective informations in a 6-week, randomized, blinded-assessment study. Psychopharmacology. 2005;178:514-523. 3. Lesem MD, Zajecka JM, Swift RH, Reeves KR, Harrigan EP. Intramuscular ziprasidone, 2 mg versus 10 mg, in the short-term management of agitated psychotic patients. J Clin Psychiatry. 2000;61:933-941.

Revised May 2005



Unique Delivery.

The **first** antidepressant patch

EMSAM* is the first and only transdermal monoamine oxidase inhibitor (MAOI) for treating depressive symptoms in patients with major depressive disorder (MDD).

Please see IMPORTANT SAFETY INFORMATION, including **Boxed WARNING**, on next page.



Unique Delivery. Proven Results.

IMPORTANT SAFETY INFORMATION

Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children
and adolescents with major depressive disorder (MDD) and other psychiatric disorders. All pediatric patients
being treated with antidepressants for any indication should be observed closely for clinical worsening,
suicidality, or unusual changes in behavior, especially during the initial few months of a course of drug therapy,
or at time of dose changes, either increases or decreases. Families and caregivers should be advised for the
need for close observation and communication with the prescriber. EMSAM is not approved for use in pediatric
patients (see Boxed WARNING)

Pooled analyses of short-term (4 to 16 weeks) placebo-controlled trials of nine antidepressant drugs (SSRIs and others) in children and adolescents with major depressive disorder (MDD), obsessive compulsive disorder (OCD), or other psychiatric disorders (a total of 24 trials involving over 4400 patients) have revealed a greater risk of adverse events representing suicidal thinking and behavior (suicidality) during the first few months of treatment in those receiving antidepressants. The average risk of such events in patients receiving antidepressants was 4%, twice the placebo risk of 2%. No suicides occurred in these trials

- To reduce the risk of hypertensive crisis, which is potentially life-threatening, foods and beverages high in tyramine must be avoided while on EMSAM 9 mg/24 hr or 12 mg/24 hr, and for 2 weeks following discontinuation of EMSAM at these doses or reducing the dose to EMSAM 6 mg/24 hr
- Due to the potential for serotonin syndrome, which is potentially life-threatening, EMSAM should not be used with
 the following antidepressants: selective serotonin reuptake inhibitors (SSRIs), dual serotonin and norepinephrine reuptake
 inhibitors (SNRIs), tricyclic antidepressants (TCAs), monoamine oxidase inhibitors (MAOIs), mirtazapine, and bupropion;
 meperidine and analgesics such as: tramadol, methadone, propoxyphene, and pentazocine; the antitussive
 dextromethorphan; cyclobenzaprine; oral selegiline; and St. John's wort
- After stopping treatment with SSRIs, SNRIs, TCAs, MAOIs, mirtazapine, bupropion; meperidine and analgesics such as: tramadol, methadone, and propoxyphene; dextromethorphan; St. John's wort; and buspirone, approximately 1 week (5 weeks for fluoxetine) should elapse before starting therapy with EMSAM. At least 2 weeks should elapse after stopping EMSAM before starting therapy with buspirone or a drug that is contraindicated with EMSAM
- Carbamazepine and oxcarbazepine are contraindicated in patients taking MAO inhibitors, including EMSAM
- The use of EMSAM is contraindicated for use with sympathomimetic amines, including amphetamines as well as cold
 products and weight-reducing preparations that contain vasoconstrictors (eg, pseudoephedrine, phenylephrine,
 phenylpropanolamine, and ephedrine)
- Patients taking EMSAM should not undergo elective surgery requiring general anesthesia or be given local anesthesia containing sympathomimetic vasoconstrictors
- EMSAM should not be used in the presence of pheochromocytoma since such tumors secrete pressor substances
- Adults with MDD or co-morbid depression in the setting of other psychiatric illness being treated with antidepressants
 should be observed for clinical worsening and suicidality, especially during the initial few months of a course of drug
 therapy, or at times of dose changes, either increases or decreases
- Risk of bipolar disorder should be ruled out prior to initiating antidepressant therapy. EMSAM is not approved for the treatment of bipolar depression
- Due to the potential for elevated blood pressure, the use of EMSAM with buspirone is not recommended
- As with other MAOIs, postural hypotension can occur with EMSAM therapy. Dose increases in the elderly should be
 made with caution and patients should be observed closely for postural changes in blood pressure throughout treatment
- EMSAM should be used with caution in patients with certain concomitant systemic illnesses that can produce altered metabolism or hemodynamic responses
- As with other psychoactive drugs, EMSAM may have the potential to impair judgment, thinking, or motor skills.
 Patients should not drive or operate hazardous machinery until they are certain EMSAM does not impair their ability to engage in such activities
- The use of alcohol is not recommended while taking EMSAM
- EMSAM should not be used in combination with tyramine-containing nutritional supplements
- EMSAM should be used in pregnancy only if the potential benefit justifies the potential risk to the fetus. Caution should be exercised when administering EMSAM to a nursing mother
- EMSAM is contraindicated in patients with known hypersensitivity to selegiline or to any component of the transdermal system
- Treatment-emergent adverse events in short-term clinical trials that occurred at a ≥2% incidence with EMSAM and for which the incidence was greater than placebo include: application site reaction (24% vs 12%), headache (18% vs 17%), insomnia (12% vs 7%), diarrhea (9% vs 7%), dry mouth (8% vs 6%), dyspepsia (4% vs 3%), rash (4% vs 2%), pharyngitis (3% vs 2%), and sinusitis (3% vs 1%)



Proven Results.

The first and only transdermal MAOI no dietary modifications at the starting and target dose of 6 mg/24 hr

Significant relief proven short-term efficacy with longer time to relapse

Demonstrated tolerability reported sexual dysfunction similar to placebo; minimal weight change

INDICATION

EMSAM is indicated for the treatment of Major Depressive Disorder (MDD).

Dose-Dependent Dietary Modifications:

To reduce the risk of hypertensive crisis, which is potentially life-threatening, foods and beverages high in tyramine must be avoided while on EMSAM* 9 mg/24 hr and 12 mg/24 hr, and for 2 weeks following discontinuation of EMSAM at these doses, or reducing the dose to EMSAM 6 mg/24 hr.

 Estimates of the incidence of sexual dysfunction cited in product labeling may underestimate actual incidence **EVSAV** 6 mg/24 hr (selegiline transdermal system)

Unique Delivery. Proven Results.

(SELEGILINE TRANSDERMAL SYSTEM)

CONTINUOUS DELIVERY FOR ONCE-DAILY APPLICATION

Brief Summary of Prescribing Information, 04/06. For complete prescribing information please consult official package circular

Suicidality in Children and Adolescents

Suicidality in Children and Adolescents Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with major depressive disorder (MDD) and other psychiatric disorders. Anyone considering the use of EMSAM (selegiline transdermal system) or any other antidepressant in a child or adolescent must balance this risk with the clinical need. Patients who are started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised for the need for close observation and communication with the prescriber. EMSAM is not approved for use in pediatric patients. (See WARNINGS and PRECAUTIONS, Perlatric IBs)

Pooled analyses of short-term (4 to 16 weeks) placebo-controlled trials of nine antidepressant drugs Probled analyses of stort-term (4 to 16 weeks) pracesor-controlled trials of finite antidepressant original (SSRIs and others) in children and adolescents with major depressive disorder (MDD), obsessive compulsive disorder (OCD), or other psychiatric disorders (a total of 24 trials involving over 4400 patients) have revealed a greater risk of adverse events representing suicidal thinking or behavior (suicidality) during the first few months of treatment in those receiving antidepressants. The average risk of such events in patients receiving antidepressants was 4%, twice the placebo risk of 2%. No suicides occurred in these trials.

CONTRAINDICATIONS
EMSAM is contraindicated in patients with known hypersensitivity to selegiline or to any component of the transdermal system

transdermal system.

EMSAM is contraindicated with selective serotonin reuptake inhibitors (SSRIs, e.g., fluoxetine, sertraline, and paroxetine); dual serotonin and norepinephrine reuptake inhibitors (SNRIs, e.g., venlataxine and duloxetine); tricyclic antidepressants (TCAs, e.g., impramine and antitriptyline); bupropion hydrochloride: meperidine and analesic agents such as tramadol, methadone and propoxyphene; the antitussive agent dextromethorphain; St. John's wort; milratazpine; and cyclobenzaprine. EMSAM should not be used with oral selegiline or other MAO inhibitors (MAOIs e.g., isocarboxazio, phenelizine, and tranyloypromine) (see WARNINGS).

Carbamazepine and oxcarbazepine are contraindicated in patients taking selegiline (see PRECAUTIONS, Drug Interactions).

As with other MAOIs, **EMSAM** is contraindicated for use with sympathomimetic amines, including amphetamines as

As with other MAOIs, EMSAM is contrainticated or use win sympathorimetic animes, including amplications well as cold products and weight-reducing preparations that contain vasoconstrictors (e.g., pseudoephedrine, henylephrine, phenylpropanolamine, and ephedrine).

As with other MAOIs, patients taking EMSAM should not undergo elective surgery requiring general anesthesia. Also, they should not be given cocaine or local anesthesia containing sympathorimmetic vasoconstrictors. EMSAM should be discortinued at least 10 days prior to elective surgery, if surgery is necessary sooner, benzodiazepines, mivacurium, repacuronium, fentanyl, morphine, and codeline may be used cautiously.

As with other MAOIs, EMSAM is contraindicated for use in patients with pheochromocytoma.

As with other Malois, Embamic contraindicated for use in patients with precinondicytimal. Embamic is an irreversible MAO inhibitor, As a class, these compounds have been associated with hypertensive crises caused by the ingestion of foods containing high amounts of fyramine. In its entirety, the data for EMSAM 6 mg/24 hours support the recommendation that a modified diet is not required at this dose. Due to the more limited data available for EMSAM 9 mg/24 hours and 12 mg/24 hours entering these doses should follow <u>Dietary Modifications</u> Required for Patients Taking EMSAM 9 mg/24 hours and 12 mg/24 hours. (See WARNINGS and PRECAUTIONS, Drug Interactions, <u>Tyramine.</u>)

WARNINGS

Clinical Worsening and Suicide Risk

Clinical Worsening and Suicide Risk
Patients with major depressive disorder (MDD), both adult and pediatric, may experience worsening of their depression and/or the emergence of suicidal ideation and behavior (suicidality) or unusual changes in behavior, whether or not they are taking antidepressant medications, and this risk may persist until significant remission occurs. There has been a long-standing concern that antidepressants may have a role in inducing worsening of depression and the emergence of suicidality in certain patients. Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with Major Depressive Disorder (MDD) and other psychiatric disorders. Pooled analyses of short-term placebo-controlled trials of nine antidepressant drugs (SSRIs and others) in children and adolescents with MDO, OCD, or other psychiatric disorders (a total of 24 trials involving over 4400 patients) have revealed a greater risk of adverse events representing suicidal behavior or thinking (suicidality) during the first few months of treatment in those receiving antidepressants. The average risk of such events in patients receiving antidepressants was 4%, wince the placebor risk of 2%. There was considerable variation in risk among drugs, but a tendency toward an increase for almost all drugs studied. The risk of suicidality was most consistently observed in the MDD trials, but there were signals of risk arising from trials in other psychiatric indications (observed in the MDD trials, but there were signals of risk arising from trials in other psychiatric indications (observed in the suicidality risk in pediatric patients extends to longer-term use, i.e., beyond several months. It is also unknown whether the suicidality risk in pediatric patients extends to longer-term use, i.e., beyond several months. It is also unknown whether the the suicidality risk extends to adults.

sucidality risk in pediatric patients extends to longer-term use, i.e., beyond several months. It is also unknown whether the suicidality risk extends to adults.

All pediatric patients being treated with antidepressants for any indication should be observed closely for clinical worsening, suicidality, and unusual changes in behavior, especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases. Such observation would generally include at least weekly face-to-face contact with patients or their family members or caregivers during the first 4 weeks of treatment, then every other week visits for the next 4 weeks, then at 12 weeks, and as clinically indicated beyond 12 weeks. Additional contact by telephone may be appropriate between face-to-face visits.

Adults with MDD or co-morbid depression in the setting of other psychiatric illness being treated with antidepressants should be observed similarly for clinical worsening and suicidality, especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases. The following symptoms, anxiety, agitation, panic attacks, insomnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomania, and mania, have been reported in adult and pediatric patients being treated with antidepressants for major depressive disorder as well as for other indications, both psychiatric and nonpsychiatric. Although a causal link between the emergence of such symptoms and either the worsening of depression and/or the emergence of suicidality musicidality. Consideration should be given to changing the therapeutic regimen, including possibly discontinuing the medication, in patients whose depression is persistently worse, or who are experiencing emergent suicidality or symptoms that might be precursors to worsening depression or suicidality, especially if these symptoms are severe, abrupt in onset, or were not part of the pati

If the decision has been made to discontinue treatment, medication should be tapered, as rapidly as is feasible, but with recognition that abruy discontinuation can be associated with certain symptoms.

Families and caregivers of pediatric patients being treated with antidepressants for major depressive disorder or other indications, both psychiatric and nonpsychiatric, should be alerted about the need to monitor patients for the emergence of agitation, irritability, unusual changes in behavior, and the other symptoms described above, as well as the emergence of suicidality, and to report such symptoms immediately to healthcare providers. Such monitoring should include daily observation by families and caregivers. Prescriptions for EMSAM should be written for the smallest quantity consistent with good patient management, in order to reduce the risk of overdose. Families and caregivers of adults being treated for depression should be similarly advised.

Screening Patients for Bipolar Disorder

Screening Patients for Bipolar Disorder
A major depressive episode may be the initial presentation of bipolar disorder. It is generally believed (though not established in controlled trials) that treating such an episode with an antidepressant alone may increase the likelihood of precipitation of a mixeo/mario episode in patients at risk for bipolar disorder. Whether any of the symptoms described above represent such a conversion is unknown. However, prior to initiating treatment with an antidepressant, patients with depressive symptoms should be adequately screened to determine if they are at risk for bipolar disorder, such screening should include a detailed psychiatric history, including a family history of suicide, bipolar disorder, and depression, it should be noted that EMSAM is not approved for use in treating bipolar depression.

Hypertensive Crisis

EMSAM is an irreversible MAO inhibitor. MAO is important in the catabolism of dietary amines (e.g., tyramine). In this regard, significant inhibition of intestinal MAO-A activity can impose a cardiovascular safety risk following the ingestion of tyramine-rich foods. As a class, MAOIs have been associated with hypertensive crises caused by the ingestion of

of tyramine-rich foods. As a class, MAOIs have been associated with hypertensive crises caused by the ingestion of foods with a high concentration of tyramine.

Hypertensive crises, which in some cases may be fatal, are characterized by some or all of the following symptoms: occipital headache which may radiate frontally, palpitation, neck stiffness or soreness, nausea, vomiting, sweating sometimes with cold, clammy skin), dilated pupils, and photophobia. Either tachycardia or bradycardia may be present and can be associated with constricting chest pain. Intracranial bleeding has been reported in association with the increase in blood pressure. Patients should be instructed as to the signs and symptoms of severe hypertension and advised to seek immediate medical attention if these signs or symptoms are present. In 6 of the 7 clinical studies conducted with EMSAM at doses of 6 mg/24 hours—12 mg/24 hours, patients were not limited to a modified diet typically associated with this class of compounds. Although no hypertensive crises were reported as part

of the safety assessment, the likelihood of developing this reaction cannot be fully determined since the amount of tyramine typically consumed during the course of treatment is not known and blood pressure was not continuously monitored. To further define the likelihood of hypertensive crises with use of EMSAM (selegiline transdermal system), severe hasse I tyramine challenge studies were conducted both with and without food (see PRECAUTIONS, Drug Interactions, Internation). In its entirety, the data for EMSAM 6 mg/24 hours support the recommendation that a modified diet is not required at this dose. Due to the more limited data available for EMSAM 9 mg/24 hours, and the results from the Phase I tyramine challenge study in fed volunteers administered EMSAM 12 mg/24 hours and 12 mg/24 hours. Brequired for Patients Taking EMSAM 9 mg/24 hours and 12 mg/24 hours.

If a hypertensive crisis occurs, EMSAM should be discontinued immediately and therapy to lower blood pressure should be instituted immediately. Phentolamine 5 mg or labetalol 20 mg administered slowly introduced should be managed by means of external cooling. Patients must be closely monitored until symptoms have stabilized.

symptoms have stabilized.

Dietary Modifications Required for Patients Taking EMSAM 9 mg/24 hours and 12 mg/24 hours

The following foods and beverages should be avoided beginning on the first day of EMSAM 9 mg/24 hours or 12 mg/24 hours treatment and should continue to be avoided for 2 weeks after a dose reduction to EMSAM 6 mg/24 hours or following the discontinuation of EMSAM 9 mg/24 hours or 12 mg/24 hours.

Food and beverages to avoid and those which are acceptable1:

Class of Food and Beverage	Tyramine-Rich Foods and Beverages to Avoid	Acceptable Foods, Containing No or Little Tyramine
Meat, Poultry and Fish	Air dried, aged and fermented meats, sausages and salamis (including cacciatore, hard salami and mortadells); pickled herring; and any spolled or improperly stored meat, poultry and fish (e.g., foods that have undergone changes in coloration, odor, or become moldy); spolled or improperly stored animal livers	Fresh meat, poultry and fish, including fresh processed meats (e.g., lunch meats, hot dogs, breakfast sausage, and cooked sliced ham)
<u>Vegetables</u>	Broad bean pods (fava bean pods)	All other vegetables
Dairy	Aged cheeses	Processed cheeses, mozzarella, ricotta cheese, cottage cheese and yogurt
Beverages	All varieties of tap beer and beers that have not been pasteurized so as to allow for ongoing fermentation	As with other antidepressants, concomitant use of alcohol with EMSAM is not recommended (Bottled and canned beers and wines contain little or no tyramine.)
Miscellaneous	Concentrated yeast extract (e.g., Marmite), sauerkraut, most soybean products (including soy sauce and tofu), OTC supplements containing tyramine	Brewer's yeast, baker's yeast, soy milk, commercial chain-restaurant pizzas prepared with cheeses low in tyramine

Adapted from K. I. Shulman, S. E. Walker. Psychiatric Annals. 2001; 31:378-384.

Naugheoritorin, Stillmarti, S. valider, resystematic Annaes, 2001; 31:376-304.

*Usa With Other Drugs Affecting Monoamine Activity
Serious, sometimes fatal, central nervous system (CNS) toxicity referred to as the "serotonin syndrome" has been reported with the combination of non-selective MAOIs with certain other drugs, including tricyclic or selective serotonin reputake inhibitor antidepressants, amphetamines, meperdine, or pentazocine. Serotonin syndrome is characterized by signs and symptoms that may include hyperthermia, rigidity, myoclonus, autonomic instability with rapid fluctuations of the vital signs, and mental status changes that include extreme aglitation progressing to drimm and coma. Similar less severe syndromes have been reported in a few patients receiving a combination of oral selegiline with near of these agents.

coma. Similar less severe syndromes have been reported in a few patients receiving a combination of oral selegiline with one of these agents. Therefore, EMSAM should not be used in combination with selective serotonin reuptake inhibitors (SNRIs, e.g., fluoxetine, sertraline, paroxetine); dual serotonin and norepinephrine reuptake inhibitors (SNRIs, e.g., veniataxine and duloxetine); tricyclic antidepressants (TCAs, e.g., imipramine and amitriptyline); oral selegiline or other MAOIs (e.g., isocarboxazid, phenetzine, and tranyloypromine); miritazapine; bupropion hydrochloride; meperidine and analgesic agents such as tramadol, methadone, and propoxyphene; the antitussive agent dextromethorphan; or St. John's wort because of the risk of life-threatening adverse reactions. Also, EMSAM should not be used with sympathonimited amines, including amphetamines as well as cold products and weight-reducing preparations that contain vasoconstrictors (e.g., pseudoephedrine, phenyleprine, phenylpropanolamine, and ephedrine). (See CONTRAINDICATIONS.)

Concomitant use of EMSAM with buspirone hydrochloride is not advised since several cases of elevated blood pressure have been reported in patients taking MAOIs who were then given buspirone HCI. After stopping treatment with SSRIs; STNIs; TCAs; MAOIs, meperidine and analgesics such as tramadol, methadone, and propoxyphene; dextromethorphan; St. John's wort; mitazapine; bupropion HCI; or buspirone HCI, a time period equal to 4-5 half-lines (approximately 1 week) of the drug or any active metabolite should elapse before starting therapy with EMSAM. Because of the long half-life of fluoxetine and its active metabolite, at least 5 weeks should elapse between discontinuation of fluoxetine and initiation of treatment with EMSAM. At least 2 weeks should elapse testing therapy with buspirone HCI or a drug that is contraindicated with EMSAM.

PRECAUTIONS

General

Hypotension: As with other MAOIs, postural hypotension, sometimes with orthostatic symptoms, can occur with

EMSAM therapy. In short-term, placebo-controlled depression studies, the incidence of orthostatic hypotension (i.e.,
a decrease of 10 mmHg or greater in mean blood pressure when changing position from supine or sitting to standing)
was 9.8% in EMSAM-treated patients and 6.7% in placebo-reated patients. It is recommended that eldry patients
treated with EMSAM be closely observed for postural changes in blood pressure throughout treatment. Dose increases
should be made cautiously in patients with pre-existing orthostasis. Postural hypotension may be relieved by having
the patient recline until the symptoms have abated. Patients should be cautioned to change positions gradually.
Patients displaying orthostatic symptoms should have appropriate dosage adjustments as warranted.

Activation of Mania/Hypomania: During Phase Iil trials, a manic reaction occurred in 8/2036 (0.4%) patients treated with EMSAM. Activation of mania/hypomania can occur in a small proportion of patients with major affective disorder treated with other marketed antidepressants. As with all antidepressants, EMSAM should be used cautiously in patients with a history of mania.

<u>Use in Patients With Concomitant Illness</u>: Clinical experience with EMSAM in patients with certain concomitant systemic illnesses is limited. Caution is advised when using EMSAM in patients with disorders or conditions that can produce altered metabolism or hemodynamic responses. EMSAM has not been systematically evaluated in patients with a history of recent myocardial infarction or unstable heart disease. Such patients were generally evaluated from clinical studies during the product's premarketing testing. No ECG abnormalities attributable to EMSAM were observed in clinical trials.

Although studies of phenylpropanolamine and pseudoephedrine did not reveal pharmacokinetic drug interactions with EMSAM, it is prudent to avoid the concomitant use of sympathomimetic agents, such as some decongestants.

Information for Patients

Information for Patients
Prescribers or other health professionals should inform patients, their families, and their caregivers about the benefits and risks associated with treatment with EMSAM and should counsel them in its appropriate use. A patient Medication Guide About Using Antidepressants in Children and Teenagers is available for EMSAM. The prescriber or health professional should instruct patients, their families, and their caregivers to read the Medication Guide and should assist them in understanding its contents. Patients should be given the opportunity to discuss the contents of the Medication Guide and to obtain answers to any questions they may have. The complete text of the Medication Guide is reprinted at the end of this document.

Patients should be advised of the following issues and asked to alert their prescriber if these occur while taking EMSAM.

Clinical Worsenia and Suicide Risk

Patients, their families and their caregivers should be encouraged to be alert to the emergence of anxiety, agitation, panic attacks, rissonnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomatia, mania, other unusual changes in behavior, worsening of depression, and suicidal ideation, especially early during antidepressant treatment or when the dose is adjusted up or down. Families and caregivers of patients should be advised to observe for the emergence of such symptoms on a day-to-day basis, since changes may be abrupt. Such symptoms should be reported to the patient's prescriber or health professional, especially if they are severe, abrupt in onset, or were not part of the patient's presenting symptoms. Symptoms such as these may be associated with an increased risk for excited this contribution and nostify change in the medication. suicidal thinking and behavior and indicate a need for very close monitoring and possibly change in the medication.

General Patients should be advised not to use oral selegiline while on EMSAM therapy. Patients should be advised not to use carbamazepine or oxcarbazepine while on EMSAM therapy. Patients should be advised not to use meperidine and analgesic agents such as tramadol, methadone, and propoxyphene. Patients should be advised not to use sympathomimetic agents while on EMSAM therapy.

Patients should be advised not to use selective serotonin reuntake inhibitors (SSRIs, e.g., fluovetine, sertraline Patients should be advised not to use selective serotonin reuptake inhibitors (SSRIs, e.g., fluoxetine, sertraine) paroxetine, and St. John's wort, dual serotonin and norepinephrine reuptake inhibitors (SNRIs, e.g., venlafaxine and duloxetine), tricyclic antidepressants (TCAs, e.g., imipramine and amitriptyline), mirtazapine, oral selegiline or other MAOIs (e.g., isocarboxazid, phenetzine, and tranylcypromline), bupropion hydrochorded or buspirone hydrochordenie while on EMSAM (selegiline transdermal system) therapy.

EMSAM has not been shown to limpair psychomotor performance; however, any psychoactive drug may potentially impair judgment, thinking, or motor skills. Patients should be cautioned about operating hazardous machinery, including automobiles, until they are reasonably certain that EMSAM therapy does not impair their ability to engage in such activities.

Patients should be told that, although EMSAM has not been shown to increase the impairment of mental and motor skills caused by alcohol, the concomitant use of EMSAM and alcohol in depressed patients is not recommended.

skills caused of succinct, the concominant use or tremSMM and actional in depressed patients is not recommended. Patients should be advised to notify their physician if they are taking, or plan to take, any prescription or over-the-counter drugs, including herbals, because of the potential for drug interactions. Patients should also be advised to avoid tyramine-containing nutritional supplements and any cough medicine containing dextromethorphan. Patients should be advised to use EMSAM exactly as prescribed. The need for dietary modifications at higher doses should be explained, and a brief description of hypertensive crisis provided. Rare hypertensive reactions with oral selegiline at doses recommended for Parkinson's disease and associated with dietary influences have been reported.

Seteginical designs of the Commence of PMSAM is unknown.

Patients should be advised that certain tyramine-rich foods and beverages should be avoided white on EMSAM 9 mg/24 hours or EMSAM 12 mg/24 hours, and for 2 weeks following discontinuation of EMSAM at these doses (see CONTRAINDICATIONS and WARNINGS).

Patients should be instructed to immediately report the occurrence of the following acute symptoms; severe

headache, neck stiffness, heart racing or palpitations, or other sudden or unusual symptoms.

Patients should be advised to avoid exposing the EMSAM application site to external sources of direct heat, such as heating pads or electric blankets, heat lamps, saunas, hot tubs, heated water beds, and prolonged direct sunlight since ay result in an increase in the amount of selegiline absorbed from the EMSAM patch and produce elevated serum levels of selegiline.

section towers or seriegimer.

Patients should be advised to change position gradually if lightheaded, faint, or dizzy while on EMSAM therapy.

Patients should be advised to notify their physician if they become pregnant or intend to become pregnant during EMSAM therapy.

Patients should be advised to notify their physician if they are breast-feeding an infant.

While patients may notice improvement with EMSAM therapy in 1 to several weeks, they should be advised of the importance of continuing drug treatment as directed.

Patients should be advised not to cut the EMSAM system into smaller portions.

For instructions on how to use EMSAM, see DOSAGE AND ADMINISTRATION, How to Use EMSAM.

The potential for drug interactions between EMSAM and a variety of drugs was examined in several human studies. Drug interaction studies described below were conducted with EMSAM 6 mg/24 hours. Although no differences are expected, drug interaction studies have not been conducted at higher doses (see <u>In vitro Metabolism</u> in Full Prescribing Information). In all of the studies described below, no drug-related adverse events were noted that required discontinuation of any subjects. Further, the incidence and nature of the adverse events were consistent with those known for seleciline or the test agent.

<u>Alcohol</u>: The pharmacokinetics and pharmacodynamics of alcohol (0.75 mg/kg) alone or in combination with **EMSAM** 6 mg/24 hours for 7 days of treatment was examined in 16 healthy volunteers. No clinically significant differences were observed in the pharmacokinetics or pharmacodynamics of alcohol or the pharmacokinetics of selegiline during co-administration. Although **EMSAM** has not been shown to increase the impairment of mental and motor skills caused by alcohol (0.75 mg/kg) and failed to alter the pharmacokinetic properties of alcohol, patients should be advised that the use of alcohol is not recommended while taking **EMSAM**.

Alprazolam: In subjects who had received EMSAM 6 mg/24 hours for 7 days, co-administration with alprazolam (15 mg/day), a CYP3A4/5 substrate, did not affect the pharmacokinetics of either selegiline or alprazolan

Carbamazepine: Carbamazepine is an enzyme inducer and typically causes decreases in drug exposure; however, slightly increased levels of selegiline and its metabolites were seen after single application of EMSAM 6 mg/24 hours in subjects who had received carbamazepine (400 mg/day) for 14 days. Changes in plasma selegiline concentrations were nearly two-fold, and variable across the subject population. The clinical relevance of these observations is unknown. Carbamazepine is contraindicated with MACIs, including selegiline (see CONTRAINDICATIONS).

Ibuprofen: In subjects who had received EMSAM 6 mg/24 hours for 11 days, combined administration with the CYP2C9 substrate ibuprofen (800 mg single dose) did not affect the pharmacokinetics of either selegiline or ibuprofen.

Ketoconazole: Seven-day treatment with ketoconazole (200 mg/day), a potent inhibitor of CYP3A4, did not affect the steady-state pharmacokinetics of selegiline in subjects who received EMSAM 6 mg/24 hours for 7 days and no differences in the pharmacokinetics of ketoconazole were observed.

<u>Levothyroxine</u>: In healthy subjects who had received **EMSAM** 6 mg/24 hours for 10 days, single dose administration with levothyroxine (150 μ g) did not alter the pharmacokinetics of either selegilline or levothyroxine (as judged by T₃ and

Olanzapine: In subjects who had received EMSAM 6 mg/24 hours for 10 days, co-administration with olanzapine, a substrate for CYP1A2, CYP2D6, and possibly CYP2A6, did not affect the pharmacokinetics of either selegiline or olanzapine.

Phenylpropanolamine (PPA): In subjects who had received EMSAM 6 mg/24 hours for 9 days, co-administration with PPA (25 mg every 4 hours for 24 hours) did not affect the pharmacokinetics of PPA. There was a higher incidence of significant blood pressure elevations with the co-administration of **EMSAM** and PPA than with PPA alone, suggesting a possible pharmacodynamic interaction. It is prudent to avoid the concomitant use of sympathomimetic agents with EMSAM.

<u>Pseudoephedrine</u>: EMSAM 6 mg/24 hours for 10 days, co-administered with pseudoephedrine (60 mg, 3 times a day) did not affect the pharmacokinetics of pseudoephedrine. The effect of pseudoephedrine on EMSAM was not examined. There were no clinically significant changes in blood pressure during pseudoephedrine administration alone, or in combination with EMSAM. Nonetheless, it is prudent to avoid the concomitant use of sympathomimetic agents with EMSAM.

Risperidone: In subjects who had received EMSAM 6 mg/24 hours for 10 days, co-administration with risperidone (2 mg per day for 7 days), a substrate for CYP2D6, did not affect the pharmacokinetics of either selegiline or risperidone.

Tyramine: Selegiline (the drug substance of EMSAM) is an irreversible inhibitor of monoamine oxidase (MAO), a

<u>Tyramine</u>: Selegiline (the drug substance of EMSAM) is an irreversible inhibitor of monoamine oxidase (MAO), a ubiquitous intracellular enzyme. MAP exists as two isoenzymes, referred to as MAO-A and MAO-B. Selegiline shows greater affinity for MAO-B; however, as selegiline concentration increases, this selectivity is lost with resulting doserelated inhibition of MAO-A; intestinal MAO is predominantly type A, while in the brain both iscenzymes exist.

MAO plays a vital physiological role in terminating the biological activity of both endogenous and exogenous amines. In addition to their role in the catabolism of monoamines in the CNS, MAOs are also important in the catabolism exogenous amines found in a variety of foods and drugs. MAO in the gastorintestinal tract (primarily type A) provides protection from exogenous amines with vasopressor actions, such as tyramine, which if absorbed intact can cause a hypertensive crisis, the so-called "cheese reaction." If a large amount of tyramine is absorbed systemically, it is taken by adrenergic neurons and causes norepinephrine release from neuronal storage sites with resultant elevation of blood pressure. While most foods contain negligible amounts or no tyramine, a few food products (see WARMINGS) may continue argue amount for tyramine is not furname that pronesent a notential risk for patients with simificant inhibition of trinestinal MAO-A resulting

pressure. Write most roous contain negigible amounts of no tyramine, a few food products (see warminus) may contain large amounts of tyramine that represent a potential risk for patients with significant inhibition of intestinal MAO-A resulting from administration of MAOIs. Tyramine-containing nutritional supplements should be avoided by patients taking EMSAM. Animal studies have indicated the transdermal administration of selegilline via EMSAM 6 mg/24 hours allows for critical levels of MAO inhibition to be achieved in the brain while avoiding levels of gastrointestinal inhibitor. To further define the risk of hypertensive crises with use of EMSAM, several Phase I tyramine challenge studies were conducted

both with and without food.

Fourteen tyramine challenge studies including 214 healthy subjects (age range 18-65; 31 subjects >50 years of age) were conducted to determine the pressor effects of oral fryamine with concurrent EMSAM treatment (6 mg/24 hours-12 mg/24 hours), measured as the dose of tyramine required to raise systolic blood pressure by 30 mmHg (TYR30). Studies were conducted with and without concomitant administration of food. Studies conducted with food are most relevant to clinical practice since tyramine typically will be consumed in food. A high-tyramine meal is considered to contain up to 40 mg of tyramine.

contain up to 40 mg or tryatmine.

One study using a crossover design in 13 subjects investigated tyramine pressor doses (TYR30) after administration of EMSAM 6 mg/24 hours and oral selegiline (5 mg twice daily) for 9 days. Mean pressor doses (TYR30) of tyramine capsules administered without food were 338 mg and 385 mg in subjects treated with EMSAM and oral selegiline, respectively. Another study using a crossover design in 10 subjects investigated tyramine pressor doses after administration of EMSAM 6 mg/24 hours or tranylcypromine 30 mg/day for 10 days. Mean pressor doses (TYR30) of tyramine capsules

administered without food were 270 mg in subjects treated with EMSAM 6 mg/24 hours and 10 mg in subjects treated with tranylcypromine.

with transportantial to the mean tyramine without food was administered to 12 subjects. The mean tyramine pressor doses (TYR30) after administration of EMSAM 6 mg/24 hours for 9 and 33 days were 292 mg and 204 mg, respectively. The lowest pressor dose was 50 mg in one subject in the 33-day group. Tyramine pressor doses were also studied in 11 subjects after extended treatment with EMSAM 12 mg/24 hours. At 30, 60, and 90 days, the mean pressor doses (TYR30) of tyramine administered without food were 95 mg, 72 mg, and 88 mg, respectively. The lowest pressor dose without food was 25 mg in 3 subjects at day 30 while on EMSAM 12 mg/24 hours.

 $Eight subjects from this study, with a mean tyramine \ pressor dose of 64 \ mg \ at 90 \ days, were subsequently administered tyramine$ with food, resulting in a mean pressor dose of 172 mg (2.7 times the mean pressor dose observed without food, p <0.003)

With the exception of one study (N=153), the Phase III clinical development program was conducted without requiring a modified diet (N=2553, 1606 at 6 mg/24 hours, and 947 at 9 mg/24 hours or 12 mg/24 hours). No hypertensive crises were reported in any patient receiving **EMSAM** (selegiline transdermal system).

In its entirety, the data for EMSAM 6 mg/24 hours support the recommendation that a modified diet is not required at this dose. Due to the more limited data available for EMSAM 9 mg/24 hours and 12 mg/24 hours, patients receiving these doses should follow Dietary Modifications Required for Patients Taking EMSAM 9 mg/24 hours and 12 mg/24 hours. (See WARNINGS.)

Warfarin: Warfarin is a substrate for CYP2C9 and CYP3A4 metabolism pathways. In healthy volunteers titrated with Coumadin® (warfarin sodium) to clinical levels of anticoagulation (INR of 1.5 to 2), co-administration with EMSAM 6 mg/24 hours for 7 days did not affect the pharmacokinetics of the individual warfarin enantiomers. EMSAM did not alter the clinical pharmacodynamic effects of warfarin as measured by INR, Factor VII or Factor X levels

Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis: In an oral carcinogenicity study in rats, selegiline given in the diet for 104 weeks was not carcinogenic up to the highest evaluable dose tested (3.5 mg/kg/day, which is 3 times the oral maximum recommended human dose on a mg/m2 basis).

Carcinogenicity studies have not been conducted with transdermal administration of selegiline

Mutagenesis: Selegiline induced mutations and chromosomal damage when tested in the in vitro mouse lymphoma assay with and without metabolic activation. Selegiline was negative in the Ames assay, the in vitro mammalian chromosome aberration assay in human lymphocytes, and the in vivo oral mouse micronucleus assay.

Impairment of Fertility: A mating and fertility study was conducted in male and female rats at transdermal doses of 10, 30, and 75 mg/kg/day of selegiline (8, 24 and 60 times the maximum recommended human dose of EMSAM [12 mg/ 24 hours] on a mg/m2 basis). Slight decreases in sperm concentration and total sperm count were observed at the high dose; however, no significant adverse effects on fertility or reproductive performance were observed.

Teratogenic Effects - Pregnancy Category C

In an embryofetal development study in rats, dams were treated with transdermal selegiline during the period of organogenesis at doses of 10, 30, and 75 mg/kg/day (8, 24, and 60 times the maximum recommended human dose [MRHD] of **EMSAM** [12 mg/24 hours] on a mg/m² basis). At the highest dose there was a decrease in fetal weight and slight increases in malformations, delayed ossification (also seen at the mid dose), and embryofetal post-implantation lethality. Concentrations of selegiline and its metabolites in fetal plasma were generally similar to those in maternal plasma. In an oral embryofetal development study in rats, a decrease in fetal weight occurred at the highest dose tested (36 mg/kg; no-effect dose 12 mg/kg); no increase in malformations was seen.

In an embryofetal development study in rabbits, dams were treated with transdermal selegiline during the period of organogenesis at doses of 2.5, 10, and 40 mg/kg/day (4, 16, and 64 times the MRHD on a mg/m² basis). A slight increase in visceral malformations was seen at the high dose. In an *oral* embryofetal development study in rabbits, increases in total resorptions and post-implantation loss, and a decrease in the number of live fetuses per dam, occurred at the highest dose tested (50 mg/kg; no-effect dose 25 mg/kg).

In a prenatal and postnatal development study in rats, dams were treated with transdermal selegiline at doses of 10, 30, and 75 mg/kg/day (8, 24, and 60 times the MRHD on a mg/m² basis) on days 6-21 of gestation and days 1-21 of the lactation period. An increase in post-implantation loss was seen at the mid and high doses, and an increase in stillborn pups was seen at the high dose. Decreases in pup weight (throughout lactation and post-weaning periods) and survival (throughout lactation period), retarded pup physical development, and pup epididymal and testicular hypoplasia, were seen at the mid and high doses. Retarded neurobehavioral and sexual development was seen at all doses. Adverse effects on pup reproductive performance, as evidenced by decreases in implantations and litter size, were seen at the high dose. These findings suggest persistent effects on the offspring of treated dams. A no-effect dose was not established for developmental toxicity. In this study concentrations of selegiline and its metabolites in milk were ~ 15 and 5 times, respectively, the concentrations in plasma, indicating that the pups were directly dosed during the lactation period.

There are no adequate and well-controlled studies in pregnant women. EMSAM should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Labor and Delivery
The effect of EMSAM on labor and delivery in humans is unknown.

In a prenatal and postnatal study of transdermal selegiline in rats, selegiline and metabolites were excreted into the milk of lactating rats. The levels of selegilline and metabolites in milk were approximately 15 and 5 times, respectively, steady-state levels of selegilline and metabolites in maternal plasma. It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised administering EMSAM to a nursing mother.

Pediatric Use

Safety and effectiveness in the pediatric population have not been established (see BOX WARNING and WARNINGS, Clinical Worsening and Suicide Risk)

Anyone considering the use of EMSAM in a child or adolescent must balance the potential risks with the clinical need.

Geriatric Use

One hundred ninety-eight (198) elderly (≥65 years of age) patients participated in clinical studies with EMSAM 6 mg/24 hours to 12 mg/24 hours. There were no overall differences in effectiveness between elderly and younger patients. In short-term, placebo-controlled depression trials, patients age 50 and older appeared to be at higher risk for rash (4.4% EMSAM versus 0% placebo) than younger patients (3.4% EMSAM versus 2.4% placebo).

ADVERSE REACTIONS

The premarketing development program for **EMSAM** included selegiline exposures in patients and/or normal subjects from two different groups of studies: 702 healthy subjects in clinical pharmacology/pharmacokinetics studies and 2036 exposures from patients in controlled and uncontrolled major depressive disorder clinical trials. The conditions and duration of treatment with EMSAM varied and included double-blind, open-label, fixed-dose, and dose titration studies of short-term and longer-term exposures. Safety was assessed by monitoring adverse events, physical examinations, vital signs, body weights, laboratory analyses, and ECGs.

Adverse events during exposure were obtained primarily by general inquiry and recorded by clinical investigators. In the tables and tabulations that follow, standard COSTART terminology has been used to classify reported adverse events. The stated frequencies of adverse events represent the proportion of individuals who experienced, at least once, a treatment-emergent adverse event of the type listed. An event was considered treatment-emergent if it occurred for the first time or worsened while receiving therapy following baseline evaluation.

Adverse Findings Observed in Short-Term, Placebo-Controlled Trials

Adverse Events Associated with Discontinuation of Treatment: Among 817 depressed patients who received EMSAM at doses of either 3 mg/24 hours (151 patients), 6 mg/24 hours (550 patients) or 6 mg/24 hours, 9 mg/24 hours, and 12 mg/24 hours (116 patients) in placebo-controlled trials of up to 8 weeks in duration, 7.1% discontinued treatment due to an adverse event as compared with 3.6% of 668 patients receiving placebo. The only adverse event associated with discontinuation, in at least 1% of EMSAM-treated patients at a rate at least twice that of placebo, was application site reaction (2% EMSAM vs. 0% placebo).

Adverse Events Occurring at an Incidence of 2% or More Among EMSAM-Treated Patients: Table 1 enumerates adverse events that occurred at an incidence of 2% or more (rounded to the nearest percent) among 817 depressed patients who received EMSAM in doses ranging from 3 to 12 mg/24 hours in placebo-controlled trials of up to 8 weeks in duration. Events included are those occurring in 2% or more of patients treated with EMSAM and for which the incidence in patients treated with EMSAM was greater than the incidence in placebo-treated patients.

Only one adverse event was associated with a reporting of at least 5% in the EMSAM group, and a rate at least twice that in the placebo group, in the pool of short-term, placebo-controlled studies: application site reactions (see Application Site Reactions, below). In one such study which utilized higher mean doses of EMSAM than that in the entire study pool, the following events met these criteria: application site reactions, insomnia, diarrhea, and pharyngitis.

These figures cannot be used to predict the incidence of adverse events in the course of usual medical practice where patient characteristics and other factors differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatments, uses, and investigators. The cited figures, however, do provide the prescribing physicians with some basis for estimating the relative contribution of drug and non-drug factors to the adverse event incidence rate in the population studied.

Table 1. Treatment-Emergent Adverse Events: Incidence in Placebo-Controlled Clinical Trials for Major essive Disorder With EMSAM (selegiline transdermal system)

Body System/Preferred Term	EMSAM (N=817)	Placebo (N=668)
	(% of Patients I	Reporting Event)
Body as a Whole Headache	18	17
Digestive Diarrhea Dyspepsia	9 4	7 3
Nervous Insomnia Dry Mouth	12 8	
Respiratory Pharyngitis Sinusitis	3 3	2 1
Skin Application Site Reaction Rash	24 4	12 2

¹ Events reported by at least 2% of patients ≥ treated with EMSAM are included, except the following events which had an incidence on placebo treatment ≥ to EMSAM: infection, nausea, dizziness, pain, abdominal pain, nervousness, back pain, asthenia, anxiety, flu syndrome, accidental injury, somnolence, rhinitis, and palpitations

Application Site Reactions: In the pool of short-term, placebo-controlled major depressive disorder studies, application site reactions (ASRs) were reported in 24% of EMSAM-treated patients and 12% of placebo-treated patients. Most ASRs were mild or moderate in severity. None were considered serious. ASRs led to dropout in 2% of EMSAM-treated patients and no

In one such study which utilized higher mean doses of **EMSAM**, ASRs were reported in 40% of **EMSAM**-treated patients and 20% of placebo-treated patients. Most of the ASIs in this study were described as crythema and most resolved spontaneously, requiring no treatment. When treatment was administered, it most commonly consisted of dermatological preparations of corticosteroids.

Male and Female Sexual Dysfunction with MAO Inhibitors: Although changes in sexual desire, sexual performance and sexual satisfaction often occur as manifestations of a psychiatric disorder, they may also be a consequence of pharmacologic treatment.

phalmacoungui traduction.

Reliable estimates of the incidence and severity of untoward experiences involving sexual desire, performance, and satisfaction are difficult to obtain, in part because patients and physicians may be reluctant to discuss them. Accordingly, estimates of the incidence or untoward sexual experience and performance cited in product belief are likely to underestimate their actual incidence. Table 2 shows that the incidence rates of sexual side effects in patients with major depressive disorder are comparable to the placebo rates in placebo-controlled trials

Table 2. Incidence of Sexual Side Effects in Placebo-Controlled Clinical Trials With EMSAM

Adverse Event	EMSAM	Placebo
	IN MA	LES ONLY
	(N=304)	(N=256)
Abnormal Ejaculation	1.0%	0.0%
Decreased Libido	0.7%	0.0%
Impotence	0.7%	0.4%
Anorgasmia	0.2%	0.0%
	IN FEM	ALES ONLY
	(N=513)	(N=412)
Decreased Libido	0.0%	0.2%

There are no adequately designed studies examining sexual dysfunction with EMSAM treatment.

Vital Sign Changes: EMSAM and placebo groups were compared with respect to (1) mean change from baseline in <u>Intal sign Changes: EMSAM</u> and piacebo groups were compared with respect to (1) mean change from baseline in vital signs (pulse, systolic blood pressure, and diastolic blood pressure) and (2) the incidence of patients meeting criteria for potentially clinically significant changes from baseline in these variables. In the pool of short-term, placebo-controlled major depressive disorder studies, 3.0% of EMSAM-treated patients and 1.5% of placebo-treated patients experienced a low systolic blood pressure, defined as a reading less than or equal to 90 mmHg with a change from baseline of at least 20 mmHg. In one study which utilized higher mean doses of EMSAM, 6.2% of EMSAM-treated patients and no placebo-treated patients experienced a low standing systolic blood pressure by these criteria. In the pool of short-term major depressive disorder trials, 9.8% of EMSAM-treated patients and 6.7% depacto-treated patients experienced a notable orthostatic change in blood pressure, defined as a decrease of at least 10 mmHg in mean blood pressure with postural change.

<u>Weight Changes</u>: In placebo-controlled studies (6 - 8 weeks), the incidence of patients who experienced ≥5% weight gain or weight loss is shown in Table 3.

Table 3. Incidence of Weight Gain and Weight Loss in Placebo-Controlled Trials With EMSAM

	-	
Weight Change	EMSAM	Placebo
	(N=757)	(N=614)
Gained ≥ 5%	2.1%	2.4%
Lost ≥ 5%	5.0%	2.8%

In these trials, the mean change in body weight among EMSAM-treated patients was -1.2 lbs compared to +0.3 lbs in placebo-treated patients

Laboratory Changes: EMSAM and placebo groups were compared with respect to (1) mean change from baseline in various serum chemistry, hematology, and urinalysis variables and (2) the incidence of patients meeting from potentially clinically significant changes from baseline in these variables. These analyses revealed no clinically important changes in laboratory test parameters associated with EMSAM.

ECG Changes: Electrocardiograms (ECGs) from EMSAM (N=817) and placebo (N=668) groups in controlled studies were compared with respect to (1) mean change from baseline in various ECG parameters and (2) the incidence of patients meeting criteria for clinically significant changes from baseline in these variables.

No clinically meaningful changes in ECG parameters from baseline to final visit were observed for patients in

controlled studies

Other Events Observed During the Premarketing Evaluation of EMSAM
During the premarketing assessment in major depressive disorder, EMSAM was administered to 2036 patients in Phase
Ill studies. The conditions and duration of exposure to EMSAM varied and included double-blind and open-label studies.
In the tabulations that follow, reported adverse events were classified using a standard COSTART—based dictionary
terminology, Ali reported adverse events are included except those already listed in Table 1 or elsewhere in labeling,
and those events occurring in only one patient. It is important to emphasize that although the events occurred during
treatment with EMSAM, they were not necessarily caused by it.
Events are further categorized by body system and listed in order of decreasing frequency according to the following
definitions: frequent adverse events are those occurring on one or more occasions in at least 1/100 patients; infrequent
adverse events are those occurring in less than 1/100 patients but at least 1/1000 patients; rare events are those
occurring in fewer than 1/1000 patients.

Body as a Wholle: Frequent: Chest pain, peck pain. Intrequent: Bacterial infection, fever, cyst, fungal infection, chills.

Body as a Whole: Fraguent: Chest pain, neck pain. Infrequent: Bacterial infection, fever, cyst, fungal infection, chills, viral infection, suicide attempt, neck rigidity, pelvic pain, photosensitivity reaction, face edema, flank pain, hernia, intentional injury, neoplasm, generalized edema, overdose. Rare: Body odor, halitosis, heat stroke, parasitic infection, malaise, moniliasis,

matiaise, monitiasis.

Cardiovascular System: Frequent: Hypertension. Infrequent: Vasodiiatation, tachycardia, migraine, syncope, atrial fibrillation, peripheral vascular disorder. Rare: Myocardial infarct.

Digestive System: Frequent: Constipation, flatulence, anorexia, gastroenteritis, vomiting. Infrequent: Increased appetite, thirst, periodontal abscess, eructation, gastritis, colitis, dysphagia, tongue edema, glossitis, increased salivation, abnormal liver function tests, melena, tongue disorder, tooth caries. Rare: Gl neoplasia, rectal hemorrhage.

Hemic and Lymphatic System: Frequent: Ecchymosis. Infrequent: Anemia, lymphadenopativ. Rare: Leukocytosis, laukonapia, patachia.

leukopenia, petechia.

Metabolic and Nutritional: Frequent: Peripheral edema. Infrequent: Hyperglycemia, increased SGPT, edema,

hypercholesteremia, increased SGOT, dehydration, alcohol intolerance, hyponatremia, increased lactic dehydrogenase. Rare: Increased alkaline phosphatase, bilirubinemia, hypoglycemic reaction. Musculoskeletal System: Frequent: Myalgia, pathological fracture. Infrequent: Arthralgia, generalized spasm, arthritis, myasthenia, arthrosis, tenosynovitis. Rare: Osteoporosis.

Nervous System: Frequent: Agitation, paresthesia, thinking abnormal, amnesia. Infrequent: Leg cramps, tremor, vertigo, hypertonia, twitching, emotional lability, confusion, manic reaction, depersonalization, hyperkinesias, hostilay myoclonus, circumoral paresthesia, hyperesthesia, increased libido, euphorit, neurosis, paranoid reaction. Rare: Ataxia.

Respiratory System: Frequent: Cough increased, bronchitis. Infrequent: Dyspinea, asthma, pneumonia, laryngismus.

Raze: Epistasis, laryngitis, yawn.

Skin and Appendages: Frequent: Pruritus, sweating, acne. Infrequent: Dry skin, maculopapular rash, contact dermatitis, urticaria, herpes simplex, alopecia, vesiculobullous rash, herpes zoster, skin hypertrophy, fungal dermatitis,

dermattits, urticaria, herpes simplex, alopecia, vesiculobullous rash, herpes zoster, skin hypertrophy, fungal dermattits, skin benign neoplasm. *Rare:* Ezezma.

Special Senses: *Frequent:* Taste perversion, tinnitus. *Infrequent:* Dry eyes, conjunctivitis, ear pain, eye pain, otitis media, parosmia. *Rare:* Mydriasis, otitis external, visual field defect.

Uriogenital System: *Frequent:* Urinary tract infection, urinary frequency, dysmenorrhea, metrorrhagia. *Infrequent:*Urinary tract infection (male), vaginitis, cystitis (female), hematuria (female), unintended pregnancy, dysuria (female), urinary urgency (male and female), vaginial moniliasis, menorrhagia, urinattion impaired (male), breast neoplasm (female), kidney calculus (female), vaginal hemorrhage, amenorrhea, breast pain, polyuria (female).

DRUG ABUSE AND DEPENDENCE

Controlled Substance Class EMSAM (selegiline transdermal system) is not a controlled substance.

Physical and Psychological Dependence

Several animal studies have assessed potential for abuse and/or dependence with chronic selegiline administration. None of these studies demonstrated a potential for selegiline abuse or dependence. EMSAM has not been systematically studied in humans for its potential for abuse, tolerance, or physical dependence.

While the clinical trials did not reveal any tendency for any drug-seeking behavior, these observations were not systematic and it is not possible to predict on the basis of this limited experience the extent to which a CNS-active drug will be misused, diverted, and/or abused once marketed. Consequently, patients should be evaluated carefully for a history of drug abuse, and such patients should be observed closely for signs of EMSAM misuse or abuse (e.g., development of tolerance, increases in dose, or drug-seeking behavior).

OVERDOSAGE

There are no specific antidotes for EMSAM. If symptoms of overdosage occur, immediately remove the EMSAM system and institute appropriate supportive therapy. For contemporary consultation on the management of poisoning or overdosage, contact the National Poison Control Center at 1-800-222-1222.

EMSAM is considered to be an irreversible MAOI at therapeutic doses and, in overdosage, is likely to cause excessive MAO-A inhibition, and may result in the signs and symptoms resembling overdosage with other non-selective, oral MAOI antidepressants (e.g., tranylcypromine [Parnate*], phenelzine [Nardil*], or isocarboxazide [Marplan*]).

Overdosage With Non-Selective MAO Inhibition

NOTE: The following is provided for reference only; it does not describe events that have actually been observed with selegiline in overdosage. No information regarding overdose by ingestion of EMSAM is available. Typical signs and symptoms associated with overdosage of non-selective MAOI antidepressants may not appear immediately. Delays of up to 12 hours between ingestion of drug and the appearance of signs may occur, and peak effects may not be observed for 24-48 hours. Since death has been reported following overdosage with MAOI agents, healthflisted with does monitoring during this period is pseceptial.

effects may not be observed for 24-48 hours. Since death has been reported following overdosage with MAUI agents, hospitalization with close monitoring during this period is essential.

Overdosage with MAOI agents is typically associated with CNS and cardiovascular toxicity. Signs and symptoms of overdosage may include, alone or in combination, any of the following: drowsiness, dizziness, faintness, irritability, hyperactivity, agitation, severe headache, halfucinations, trismus, opisthotonos, convulsions, coma, rapid and irregular pulse, hyperension, hypotension and vascular collapse, precordial pain, respiratory depression and failure, hyperpyrexia, diaphoresis, and cool, clammy skin. Type and intensity of symptoms may be related to extent of the

Injury private, unapholeas, and book calming shall prove the measures, with pharmacological intervention as appropriate. Symptoms may persist after drug washout because of the irreversible inhibitory effects of these agents on systemic MAO activity. With overdosage, in order to avoid the occurrence of hypertensive crisis ("cheese reaction"), dietary tyramine should be restricted for several weeks beyond recovery to permit regeneration of the peripheral MAO-A isoenzyme.

DOSAGE AND ADMINISTRATION

DOSAGE AND ADMINISTRATION
Initial Treatment

EMSAM should be applied to dry, intact skin on the upper torso (below the neck and above the waist), upper thigh or
the outer surface of the upper arm once every 24 hours. The recommended starting dose and target dose for EMSAM
is 6 mg/24 hours. EMSAM has been systematically evaluated and shown to be effective in a dose range of 6 mg/24 hours
to 12 mg/24 hours. However, the trials were not designed to assess if higher doses are more effective than the lowest
effective dose of 6 mg/24 hours. Based on clinical judgment, if dose increases are indicated for Individual patients,
they should occur in dose increments of 3 mg/24 hours (up to a maximum dose of 12 mg/24 hours) at intervals of no
less than 2 weeks. As with all antidepressant drugs, full antidepressant effect may be delayed.
Patients should be informed that tyramine-rich foods and beverages should be avoided beginning on the first day of
EMSAM 9 mg/24 hours of 2 mg/24 hours and should continue to be avoided beginning on the great and should continue to be avoided for 2 weeks after a dose

EMSAM 9 mg/24 hours or 12 mg/24 hours treatment and should continue to be avoided for 2 weeks after a dose reduction to EMSAM 6 mg/24 hours or following the discontinuation of EMSAM 9 mg/24 hours or 12 mg/24 hours (see WARNINGS)

Special Populations

Special repulations of the patients with mild to moderate renal or hepatic impairment. The recommended dose for elderly patients (≥65 years) is EMSAM 6 mg/24 hours daily. Dose increases, in the elderly, should be made with caution and patients should be closely observed for postural changes in blood pressure throughout treatment.

How to Use EMSAM

- EMSAM should be applied to dry, intact skin on the upper torso (below the neck and above the waist), upper thigh or the outer surface of the upper arm. A new application site should be selected with each new patch to avoid re-application to the same site on consecutive days. Patches should be applied at approximately the same
- awout re-application to the same site on can security days. Paraches should be applied at approximately the same time each day.

 2. Apply the patch to an area of skin that is not hairy, oily, irritated, broken, scarred or calloused. Do not place the patch where your clothing is tight, which could cause the patch to rub off.

 3. After you have selected the site for your patch, wash the area gently and thoroughly with soap and warm water. Rinse until all soap is removed. Dry the area with a clean dry towel.

 4. Just before you apply the patch, remove it from the pouch. Remove half of the protective backing and throw it
- away. Try not to touch the exposed side (sticky side) of the patch, because the medicine could come off on your
- fingers.

 5. Press the sticky side of the patch firmly against the skin site that was just washed and dried. Remove the second half of the protective liner and press the remaining sticky side firmly against your skin. Make sure that the patch is flat against the skin (there should be no bumps or folds in the patch) and is sticking securely. Be sure the edges are stuck to the skin surface.
- sure the edges are stuck to the skin surrace.

 After you have applied the patch, <u>wash your hands</u> thoroughly with soap and water to remove any medicine that may have gotten on them. Do not touch your eyes until after you have washed your hands.

 After 24 hours, remove the patch. Do not touch the sticky side. As soon as you have removed the patch, fold it so that the sticky side sticks to itself.

 Throw away the folded patch so that children and/or pets cannot reach it.
- Wash your hands with soap and water.
- wash you hallow with solp and water.

 If your patch falls off, apply a new patch to a new site and resume your previous schedule.

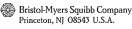
 Only one EMSAM patch should be worn at a time.

 Avoid exposing the EMSAM application site to external sources of direct heat, such as heating pads or electric
- blankets, heat lamps, saunas, hot tubs, heated water beds, and prolonged direct sunlight.

Maintenance Treatment

Maintenance Treatment
It is generally agreed that episodes of depression require several months or longer of sustained pharmacologic therapy. The benefit of maintaining depressed patients on therapy with EMSAM at a dose of 6 mg/24 hours after achieving a responder status for an average duration of about 25 days was demonstrated in a controlled trial (see Clinical Efficacy Trials in Full Prescribing Information and INDICATIONS AND USAGE). The physician who elects to use EMSAM for extended periods should periodically re-evaluate the long-term usefulness of the drug for the individual patient.

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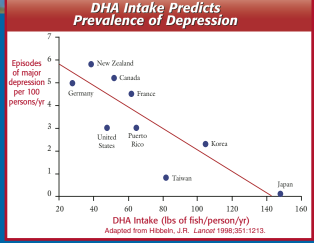


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"...reduced membrane DHA emerged as a significant predictor of depression..."

Edwards, R., et al. J Affect Disord 1998;48:149-155.

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Postpartum-"...lower DHA content in mothers' milk...[was] associated with higher rates of postpartum depression." Hibbeln, J.R. J Affect Disord 2002;69:15-29.

"Trials have shown that folate supplementation hastens recovery from depressive episodes and enhances the effect of antidepressants." Morris, M.S., et al. Psychother and Psychosom 2003;72(2):80-7.

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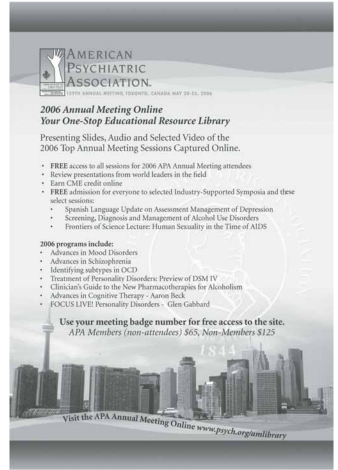
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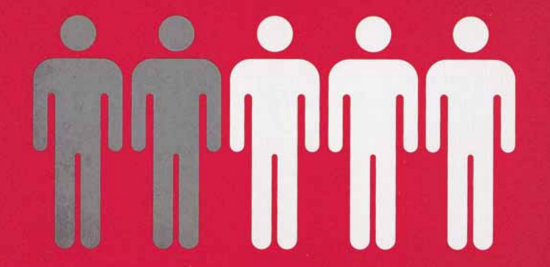
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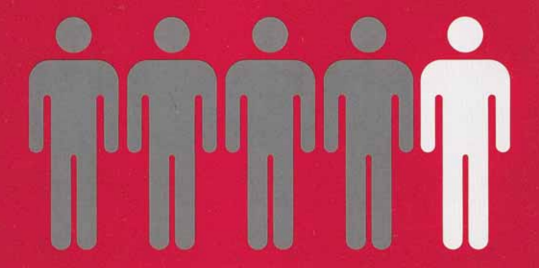


41% of all patients had the metabolic syndrome at baseline in the landmark CATIE schizophrenia study.1

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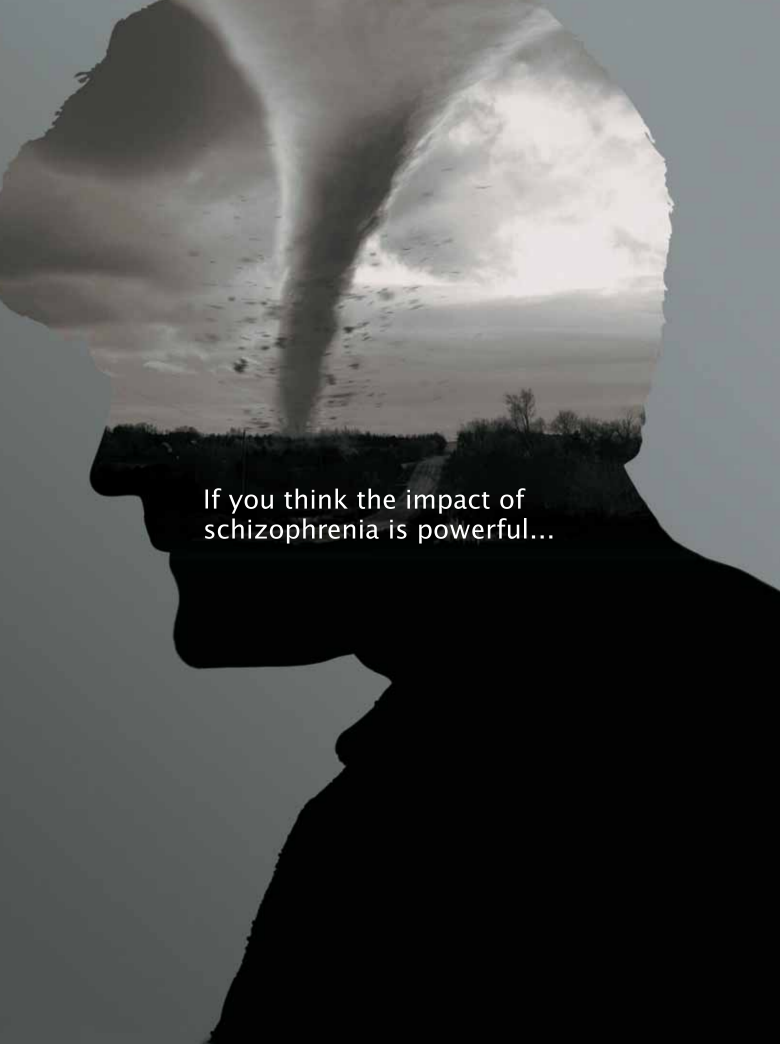


13% of patients had diabetes in the landmark CATIE schizophrenia study at baseline—4 times more common than in the general population.²

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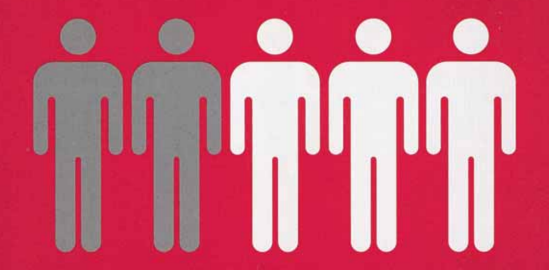
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41% of all patients had the metabolic syndrome at baseline in the landmark CATIE schizophrenia study.¹

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Mild - Moderate - Severe

ARICEPT® is indicated for the treatment of dementia of the Alzheimer's type. Efficacy has been demonstrated in patients with mild to moderate Alzheimer's disease, as well as in patients with severe Alzheimer's disease.

Cholinesterase inhibitors have the potential to increase gastric acid secretion. Patients at risk for developing ulcers, including those receiving concurrent NSAIDs, should be monitored closely for gastrointestinal bleeding.

In clinical trials, syncopal episodes have been reported (2% for ARICEPT versus 1% for placebo).

In clinical trials, the most common adverse events seen with ARICEPT were nausea, diarrhea, insomnia, vomiting, muscle cramps, fatigue, anorexia, and ecchymosis. In studies, these were mild and transient.

Please see brief summary of prescribing information on adjacent page.









Brief Summary—see package insert for full prescribing information.

ARICEPT® (Donepezil Hydrochloride Tablets)

ARICEPT® ODT (Donepezil Hydrochloride) Orally Disintegrating Tablets

INDICATIONS AND USAGE ARICEPT® is indicated for the treatment of dementia of the Alzheimer's type. Efficacy has been demonstrated in patients with mild to moderate Alzheimer's Disease, as well as in patients with severe Alzheimer's Disease. CONTRAINDICATIONS ARICEPT® is contraindicated in patients with known hypersensitivity to donepezil hydrochloride or to piperidine derivatives. WARNINGS Anesthesia: ARICEPT®, as a cholinesterase inhibitor, is likely to exaggerate succinylcholine-type muscle relaxation during anesthesia. Cardiovascular Conditions: Because of their pharmacological action, cholinesterase inhibitors may have vagolonic effects on the sinoatrial and atrioventricular nodes. This effect may manifest as bradycardia or heart block in patients both with and without known underlying cardiac conduction abnormalities. Syncopal episodes have been reported in association with the use of ARICEPT®. **Gastrointestinal Conditions:** Through their primary action, cholinesterase inhibitors may be expected to increase gastric acid secretion due to increased cholinergic activity. Therefore, patients should be monitored closely for symptoms of active or occult gastrointestinal bleeding, especially those at increased risk for developing ulcers, e.g., those with a history of ulcer disease or those receiving concurrent nonsteroidal anti-inflammatory drugs (NSAIDS). Clinical studies of ARICEPT® have shown no increase, relative to placebo, in the incidence of either peptic ulcer disease or gastrointestinal bleeding. ARICEPT®, as a predictable consequence of its pharmacological properties, has been shown to produce diarrhea, nausea and vomiting. These effects, when they occur, appear more frequently with the 10 mg/day dose than with the 5 mg/day dose. In most cases, these effects have been mild and transient, sometimes lasting one to three weeks, and have resolved during continued use of ARICEPT®.

Genitourinary: Although not observed in clinical trials of ARICEPT®, cholinomimetics may cause bladder outflow obstruction.

Neurological Conditions: Seizures: Cholinomimetics are believed to have some potential to cause generalized convulsions. However, seizure activity also may be a manifestation of Alzheimer's Disease. Pulmonary Conditions: Because of their cholinomimetic actions, cholinesterase inhibitors should be prescribed with care to patients with a history of asthma or obstructive pulmonary disease. PRECAUTIONS Drug-Drug Interactions (see Clinical Pharmacology: Clinical Pharmacokinetics: Drug-drug Interactions) Effect of ARICEPT® on the Metabolism of Other Drugs: No in vivo clinical trials have investigated the effect of ARICEPT® on the clearance of drugs metabolized by CYP 3A4 (e.g. cisapride, terfenadine) or by CYP 2D6 (e.g. imipramine). However, in vitro studies show a low rate of binding to these enzymes (mean K, about 50-130 µM), that, given the therapeutic plasma concentrations of donepezil (164 nM), indicates little likelihood of interference. Whether ARICEPT® has any potential for enzyme induction is not known. Formal pharmacokinetic studies evaluated the potential of ARICEPT® for interaction with theophylline, cimetidine, warfarin, digoxin and ketoconazole. No effects of ARICEPT® on the pharmacokinetics of these drugs were observed. Effect of Other Drugs on the Metabolism of ARICEPT®: Ketoconazole and quinidine, inhibitors of CYP450, 3A4 and 2D6, respectively, inhibit donepezil metabolism *in vitro*. Whether there is a clinical effect of quinidine is not known. In a 7-day crossover study in 18 healthy volunteers, ketoconazole (200 mg q.d.) increased mean donepezil (5 mg q.d.) concentrations (AUC₀₋₂₄ and C_{max}) by 36%. The clinical relevance of this increase in concentration is unknown. Inducers of CYP 2D6 and CYP 3A4 (e.g., phenytoin, carbamazepine, dexamethasone, rifampin, and phenobarbital) could increase the rate of elimination of ARICEPT®. Formal pharmacokinetic studies demonstrated that the metabolism of ARICEPT® is not significantly affected by concurrent administration of digoxin or cimetidine. **Use** with Anticholinergics: Because of their mechanism of action, cholinesterase inhibitors have the potential to interfere with the activity of anticholinergic medications. Use with Cholinomimetics and Other Cholinesterase Inhibitors: A synergistic effect may be expected when cholinesterase inhibitors are given concurrently with succinylcholine, similar neuromuscular blocking agents or cholinergic agonists such as bethanechol. Carcinogenesis, Mutagenesis, Impairment of Fertility No evidence of a carcinogenic potential was obtained in an 88-week carcinogenicity study of donepezil hydrochloride conducted in CD-1 mice at doses up to 180 mg/kg/day (approximately 90 times the maximum recommended human dose on a mg/m² basis), or in a 104-week carcinogenicity study in Sprague-Dawley rats at doses up to 30 mg/kg/day (approximately 30 times the maximum recommended human dose on a mg/m² basis). Donepezil was not mutagenic in the Ames reverse mutation assay in bacteria, or in a mouse lymphoma forward mutation assay in vitro. In the chromosome aberration test in cultures of Chinese hamster lung (CHL) cells, some clastogenic effects were observed. Donepezil was not clastogenic in the *in vivo* mouse micronucleus test and was not genotoxic in an *in vivo* unscheduled DNA synthesis assay in rats. Donepezil had no effect on fertility in rats at doses up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m² basis). Pregnancy Pregnancy Category C: Teratology studies conducted in pregnant rats at doses up to 16 mg/kg/day (approximately 13 times the maximum recommended human dose on a mg/m² basis) and in pregnant rabbits at doses up to 10 mg/kg/day (approximately 16 times the maximum recommended human dose on a mg/m basis) did not disclose any evidence for a teratogenic potential of donenezil. However, in a study in which pregnant rats were given up to 10 mg/kg/day (approximately 8 times the maximum recommended human dose on a mg/m² basis) from day 17 of gestation through day 20 postpartum, there was a slight increase in still births and a slight decrease in pup survival through day 4 postpartum at this dose; the next lower dose tested was 3 mg/kg/day. There are no adequate or well-controlled studies in pregnant women. ARICEPT® should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Nursing Mothers It is not known whether done pezil is excreted in human breast milk. ARICEPT® has no indication for use in nursing mothers. Pediatric Use There are no adequate and well-controlled trials to document the safety and efficacy of ARICEPT® in any illness occurring in children. Geriatric Use Alzheimer's disease is a disorder occurring primarily in individuals over 55 years of age. The mean age of the patients enrolled in the clinical studies with ARICEPT® was 73 years; 80% of these patients were between 65 and 84 years old and 49% of the patients were at or above the age of 75. The efficacy and safety data presented in the clinical trials section were obtained from these patients. There were no clinically significant differences in most adverse events reported by patient groups ≥65 years old and <65 years old. ADVERSE REACTIONS Mild To Moderate Alzheimer's Disease Adverse Events Leading to Discontinuation The rates of discontinuation from controlled clinical trials of ARICEPT® due to adverse events for the ARICEPT® 5 mg/day treatment groups were comparable to those of placebo-treatment groups at approximately 5%. The rate of discontinuation of patients who received 7-day escalations from 5 mg/day to 10 mg/day, was higher at 13%. The most common adverse events leading to discontinuation, defined as those occurring in at least 2% of patients and at twice the incidence seen in placebo patients, are shown in Table 1. Table 1. Most Frequent Adverse Events Leading to Withdrawal from Controlled Clinical Trials by Dose Group (Placebo, 5 mg/day ARICEPT*, and 10 mg/day ARICEPT*, respectively); Patients Randomized (355, 350, 315); Event/% Discontinuing: Nausea (1%, 1%, 3%); Diarrhea (0%, <1%, 3%); Vomiting (<1%, <1%, 2%). Most Frequent Adverse Clinical Events Seen in Association with the Use of ARICEPT®. The most common adverse events, defined as those occurring at a frequency of at least 5% in patients receiving 10 mg/day and twice the placebo rate, are largely predicted by ARICEPT®'s cholinomimetic effects. These include nausea, diarrhea, insomnia, vomiting, muscle cramp, fatigue and anorexia. These adverse events were often of mild intensity and transient, resolving during continued ARICEPT® treatment without the need for dose modification. There is evidence to suggest that the frequency of these common adverse events may be affected by the rate of titration. An open-label study was conducted with 269 natients who received placeho in the 15 and 30-week studies. These natients were titrated to a dose of 10 mg/day over a 6-week period. The rates of common adverse events were lower than those seen in patients titrated to 10 mg/day over one week in the controlled clinical trials and were comparable to those seen in patients on 5 mg/day. See Table 2 for a comparison of the most common adverse events following one and six week titration regimens. Table 2. Comparison of rates of adverse events in patients titrated to 10 mg/day over 1 and 6 weeks (No titration: Placebo [n=315], No titration: 5 mg/day [n=311], One week titration: 10 mg/day [n=315], Six week titration: 10 mg/day [n=269], respectively): Nausea (6%, 5% 19%, 6%); Diarrhea (5%, 8%, 15%, 9%); Insomnia (6%, 6%, 14%, 6%); Fatigue (3%, 4%, 8%, 3%); Vomiting (3%, 3%, 8%, 5%); Muscle cramps (2%, 6%, 8%, 3%); Anorexia (2%, 3%, 7%, 3%). Adverse Events Reported in Controlled Trials The events cited reflect experience gained under closely monitored conditions of clinical trials in a highly selected patient population. In actual clinical practice or in other clinical trials, these frequency estimates may not apply, as the conditions of use, reporting behavior, and the kinds of patients treated may differ. Table 3 lists treatment emergent signs and symptoms that were reported in at least 2% of patients in placebo-controlled trials who received ARICEPT® and for which the rate of occurrence was greater for ARICEPT® assigned than placebo assigned patients. In general, adverse events occurred more frequently in female patients and with advancing age. Table 3. Adverse Events Reported in Controlled Clinical Trials in Mild to Moderate Alzheimer's Disease in at Least 2% of Patients Receiving ARICEPT® and at a Higher Frequency than Placebo-treated Patients (Body System/Adverse Event: Placebo [n=355], ARICEPT® [n=747], respectively): Percent of Patients with any Adverse Event: 72, 74. Body as a Whole: Headache (9, 10); Pain, various locations (8, 9); Accident (6, 7); Fatigue (3, 5). Cardiovascular System: Syncope (1, 2). Digestive System: Nausea (6, 11); Diarrhea (5, 10); Vomiting (3, 5); Anorexia (2, 4). Hemic and Lymphatic System: Ecchymosis (3, 4). Metabolic and Nutritional Systems: Weight Decrease (1, 3). Musculoskeletal System: Muscle Cramps (2, 6); Arthritis (1, 2). Nervous System: Insomnia (6, 9); Dizziness (6, 8); Depression (<1, 3); Abnormal Draams (0, 3); Somnolence (<1, 2). Urogenital System: Frequent Urination (1, 2). Other Adverse Events Observed During Clinical Trials. ARICEPT has been administered to over 1700 individuals during clinical trials worldwide. Approximately 1200 of these patients have been treated for at least 3 months and more than 1000 patients have been treated for at least 6 months. Controlled and uncontrolled trials

in the United States included approximately 900 patients. In regards to the highest dose of 10 mg/day, this population includes 650 patients treated for 3 months, 475 patients treated for 6 months and 116 patients treated for over 1 year. The range of patient exposure is from 1 to 1214 days. Treatment emergent signs and symptoms that occurred during 3 controlled clinical trials and two open-label trials in the United States were recorded as adverse events by the clinical investigators using terminology of their own choosing. To provide an overall estimate of the proportion of individuals having similar types of events, the events were grouped into a smaller number of standardized categories using a modified COSTART dictionary and event frequencies were calculated across all studies. These categories are used in the listing below. The frequencies represent the proportion of 900 patients from these trials who experienced that event while receiving ARICEPT®. All adverse events occurring at least twice are included, except for those already listed in Tables 2 or 3, COSTART terms too general to be informative, or events less likely to be drug caused. Events are classified by body system and listed using the following definitions: frequent adverse events—those occurring in at least 1/100 patients; infrequent adverse events those occurring in 1/100 to 1/1000 patients. These adverse events are not necessarily related to ARICEPT® treatment and in most cases were observed at a similar frequency in placebo-treated patients in the controlled studies. No important additional adverse events were seen in studies conducted outside the United States. Body as a Whole: Frequent: influenza, chest pain, toothache; Infrequent: fever, edema face, periorbital edema, hernia hiatal, abscess, cellulitis, chills, generalized coldness, head fullness, listlessness. **Cardiovascular System:** Frequent: hypertension, vasodilation, atrial fibrillation, hot flashes, hypotension; Infrequent: angina pectoris, postural hypotension, myocardial infarction, AV block (first degree), congestive heart failure, arteritis, bradycardia, peripheral vascular disease, supraventricular tachycardia, deep vein thrombosis. Digestive System: Frequent: fecal incontinence, gastrointestinal bleeding, bloating, epigastric pain; Infrequent: eructation, gingivitis, increased appetite, flatulence, periodontal abscess, cholelithiasis, diverticulitis, drooling, dry mouth, fever sore, gastritis, irritable colon, tongue edema, epigastric distress, gastroenteritis, increased transaminases, hemorrhoids, ileus, increased thirst, jaundioe, melena, polydipsia, duodenal ulcer, stomach ulcer. **Endocrine** System: Infrequent: diabetes mellitus, goiter. Hemic and Lymphatic System: Infrequent: anemia, thrombocythemia, thrombocytopenia, eosinophilia, erythrocytopenia. Metabolic and Nutritional Disorders: Frequent: dehydration; Infrequent gout, hypokalemia, increased creatine kinase, hyperglycemia, weight increase, increased lactate dehydrogenase. Musculoskeletal System: Frequent: bone fracture; Infrequent: muscle weakness, muscle fasciculation. Nervous System: Frequent: delusions, tremor, irritability, paresthesia, aggression, vertigo, ataxia, increased libido, restlessness, abnormal crying, nervousness, aphasia; Infrequent: cerebrovascular accident, intracranial hemorrhage, transient ischemic attack, emotional lability, neuralgia, coldness (localized), muscle spasm, dysphoria, gait abnormality, hypertonia, hypokinesia, neurodermatitis, numbness (localized), paranoia, dysarthria, dysphasia, hostility, decreased libido, melancholia, emotional withdrawal, nystagmus, pacing. Respiratory System: Frequent: dyspnea, sore throat, bronchitis; Infrequent: epistaxis, post nasal drip, pneumonia, hyperventilation, pulmonary congestion, wheezing, hypoxia, pharyngitis, pleurisy, pulmonary collapse, sleep apnea, snoring. Skin and Appendages: Frequent: pruritus, diaphoresis, urticaria; Infrequent: dermatitis, erythema, skin discoloration, hyperkeratosis, alopecia, fungal dermatitis, herpes zoster, hirsutism, skin striae, night sweats, skin ulcer. Special Senses: Frequent: cataract, eye irritation, vision blurred; Infrequent: dry eyes, glaucoma, earache, tinnitus, blepharitis, decreased hearing, retinal hemorrhage, otitis externa, otitis media, bad taste, conjunctival hemorrhage, ear buzzing, motion sickness, spots before eyes. **Urogenital System:** Frequent: urinary incontinence, nocturia; Infrequent: dysuria, hematuria, urinary urgency, metrorrhagia, cystitis, enuresis, prostate hypertrophy, pyelonephritis, inability to empty bladder, breast fibroadenosis, fibrocystic breast, mastitis, pyuria, renal failure, vaginitis. Severe Alzheimer's Disease Adverse Events Leading to Discontinuation: The rates of discontinuation from controlled clinical trials of ARICEPT® due to adverse events for the ARICEPT® patients were approximately 12% compared to 7% for placebo patients. The most common adverse events leading to discontinuation, defined as those occurring in at least 2% of ARICEPT® patients and at twice the incidence seen in placebo patients, were anorexia (2% vs 1% placebo), nausea (2% vs <1% placebo), diarrhea (2% vs 0% placebo), and urinary tract infection (2% vs 1% placebo). Most Frequent Adverse Clinical Events Seen in Association with the Use of ARICEPT The most common adverse events, defined as those occurring at a frequency of at least 5% in patients receiving ARICEPT® and twice the placebo rate, are largely predicted by ARICEPT®'s cholinomimetic effects. These include diarrhea, anorexia, vomiting, nausea, and ecchymosis. These adverse events were often of mild intensity and transient, resolving during continued ARICEPT® treatment without the need for dose modification. Adverse Events Reported in Controlled Trials Table 4 lists treatment emergent signs and symptoms that were reported in at least 2% of patients in placebo-controlled trials who received ARICEPT® and for which the rate of occurrence was greater for ARICEPT® assigned than placebo assigned patients. Table 4. Adverse Events Reported in $\textbf{Controlled Clinical Trials in Severe Alzheimer's Disease in at Least 2\% of Patients \, \textbf{Receiving ARICEPT}^{\circ} \, \textbf{and at any the property of the Controlled Clinical Trials in Severe Alzheimer's Disease in at Least 2\% of Patients \, \textbf{Receiving ARICEPT}^{\circ} \, \textbf{and at any the Controlled Clinical Trials in Severe Alzheimer's Disease in at Least 2\% of Patients \, \textbf{Receiving ARICEPT}^{\circ} \, \textbf{and at any the Controlled Clinical Trials}^{\circ} \, \textbf{ARICEPT}^{\circ} \, \textbf{and at any the Controlled Clinical Trials}^{\circ} \, \textbf{ARICEPT}^{\circ} \, \textbf{and at any the Controlled Clinical Trials}^{\circ} \, \textbf{ARICEPT}^{\circ} \, \textbf{AR$ Higher Frequency than Placebo-treated Patients (Body System/Adverse Event: Placebo [n=392], ARICEPT[®] [n=501], respectively): Percent of Patients with any Adverse Event: 73, 81. Body as a Whole: Accident (12, 13); Infection (9, 11); Headache (3, 4); Pain (2, 3); Back Pain (2, 3); Fever (1, 2); Chest Pain (<1, 2). Cardiovascular System: Hypertension (2, 3); Hemorrhage (1, 2); Syncope (1, 2). Digestive System: Diarrhea (4, 10); Vomiting (4, 8); Anorexia (4, 8); Nausea (2, 6), Hemic and Lymphatic System: Ecchymosis (2, 5), Metabolic and Nutritional Systems: Creatine Phosphokinase Increased (1, 3); Dehydration (1, 2); Hyperlipemia (<1, 2), Nervous System: Insomnia (4, 5); Hostility (2, 3); Nervousness (2, 3); Hallucinations (1, 3); Somnolence (1, 2); Dizziness (1, 2); Depression (1, 2); Confusion (1, 2); Emotional Lability (1, 2); Personality Disorder (1, 2). Skin and Appendages: Eczema (2, 3). Urogenital System: Urinary Incontinence (1, 2). Other Adverse Events Observed During Clinical Trials ARICEPT® has been administered to over 600 patients with severe Alzheimer's Disease during clinical trials of at least 6 months duration, including 3 double blind placebo controlled trials, one of which had an open label extension. All adverse events occurring at least twice are included, except for those already listed in Table 4, COSTART terms too general to be informative, or events less likely to be drug caused. Events are classified by body system using the COSTART dictionary and listed using the following definitions: frequent adverse events—those occurring in at least 1/100 patients; infrequent adverse events—those occurring in 1/100 to 1/1000 patients. These adverse events are not necessarily related to ARICEPT® treatment and in most cases were observed at a similar frequency in placebo-treated patients in the controlled studies. Body as a Whole: Frequent: abdominal pain, asthenia, fungal infection, flu syndrome; Infrequent: allergic reaction, cellulitis, malaise, sepsis, face edema, hernia. Cardiovascular System: Frequent: hypotension, bradycardia, ECG abnormal, heart failure; Infrequent: myocardial infarction, angina pectoris, atrial fibrillation, congestive heart failure, peripheral vascular disorder, supraventricular extrasystoles, ventricular extrasystoles, cardiomegaly. Digestive System: Frequent: constipation, gastroenteritis, fecal incontinence, dyspepsia; Infrequent: gamma glutamyl transpeptidase increase, gastritis, dysphagia, periodontitis, stomach ulcer, periodontal abscess, flatulence, liver gamina gudarnyn danspeptudase increase, gasunus, dysphagda, perdodunus, stornach duce, perdodunua dassess, induderte, in function lests shormal, eructation, esophagitis, redal hemorrhage. Endocrine System: Infraquent diabetes mellitus. Hemie and Lymphatic System: Frequent: anemia; Infraquent: leukocytosis. Metabolic and Nutritional Disorders: Frequent: weight loss, peripheral edema, edema, lactic dehydrogenase increased, alkaline phosphatase increased; Infrequent: hypercholesteremia, hypokalemia, hypoglycemia, weight gain, bilirubinemia, BUN increased, B₁₂ deficiency anemia, cachexia, creatinine increased, gout, hyponatremia, hypoproteinemia, iron deficiency anemia, SGOT increased, SGPT increased. Musculoskeletal System: Frequent: arthritis; Infrequent: arthrosis, bone fracture, arthralgia, leg cramps, osteoporosis, myalgia. Nervous System: Frequent: aqitation. anxiety, tremor, convulsion, wandering, abnormal gait; *Infrequent:* apathy, vertigo, delusions, abnormal dreams, cerebrovascular accident, increased salivation, ataxia, euphoria, vasodilatation, cerebral hemorrhage, cerebral infarction, cerebral ischemia, dementia, extrapyramidal syndrome, grand mal convulsion, hemiplegia, hypertonia, hypokinesia. Respiratory System: Frequent: pharyngitis, pneumonia, cough increased, bronchitis; Infrequent: dyspnea, rhinitis, asthma. Skin and Appendages: Frequent: rash, skin ulcer, pruritus; Infrequent: psoriasis, skin discoloration, herpes zoster, dry skin, sweating, urticaria, vesiculobullous rash. Special Senses: Infrequent: conjunctivitis, glaucoma, abnormal vision, ear pain, lacrimation disorder. Urogenital System: Frequent: urinary tract infection, cystitis, hematuria, glycosuria; Infrequent: vaginitis, dysuria, urinary frequency, albuminuria. Postintroduction Reports Voluntary reports of adverse events temporally associated with ARICEPT® that have been received since market introduction that are not listed above, and that there is inadequate data to determine the causal relationship with the drug include the following: abdominal pain. agitation. cholecystitis. confusion, convulsions, hallucinations, heart block (all types), hemolytic anemia, hepatitis, hyponatremia, neuroleptic malignant syndrome, pancreatitis, and rash. **OVERDOSAGE Because strategies for the management of** overdose are continually evolving, it is advisable to contact a Poison Control Center to determine the latest recommendations for the management of an overdose of any drug. As in any case of overdose, general supportive $measures should be {\it utilized}. Overdosage {\it with cholinesterase inhibitors can result in cholinergic crisis characterized by severe nauseau and the contraction of the contraction$ vomiting, salivation, sweating, bradycardia, hypotension, respiratory depression, collapse and convulsions. Increasing muscle weakness is a possibility and may result in death if respiratory muscles are involved. Tertiary anticholinergics such as atropine may be used as an antidote for ARICEPT® overdosage. Intravenous atropine sulfate titrated to effect is recommended: an initial dose of 1.0 to 2.0 mg IV with subsequent doses based upon clinical response. Atypical responses in blood pressure and heart rate have been reported with other cholinomimetics when co-administered with quaternary anticholinergics such as glycopyrrolate. It is not known whether ARICEPT® and/or its metabolites can be removed by dialysis (hemodialysis, peritoneal dialysis, or hemofiltration). Dose-related signs of toxicity in animals included reduced spontaneous movement, prone position, staggering gait, lacrimation, clonic convulsions, essed respiration, salivation, miosis, tremors, fasciculation and lower body surface temperature

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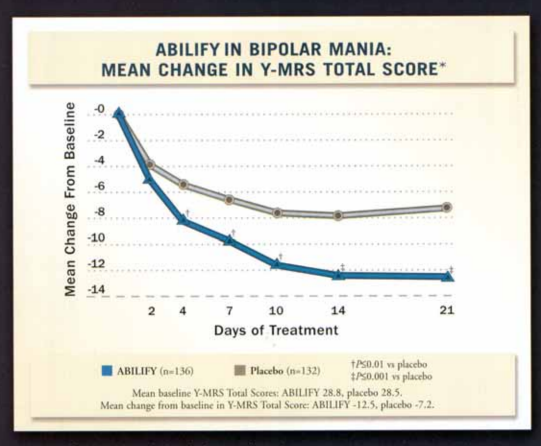
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ABILIFY® (aripiprazole) efficacy looks good on paper...



Adapted from Sachs et al. J Psychopharmacol. 2006.

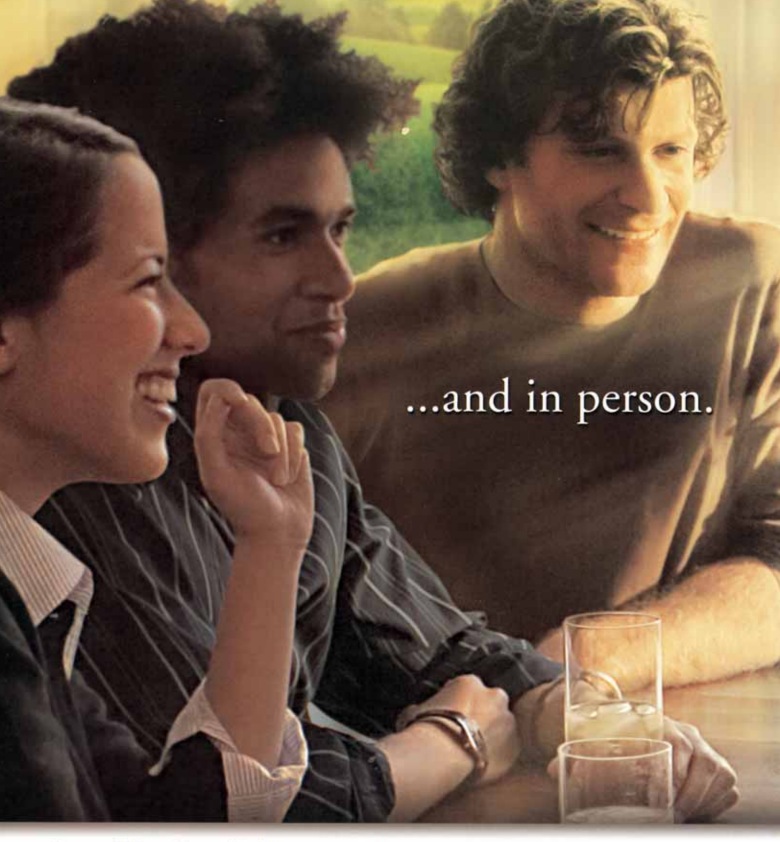
Data from a 3-week, double-blind, randomized, placebo-controlled trial in patients with Bipolar I Disorder experiencing acute manic or mixed episodes. Patients were randomized to receive either placebo or aripiprazole with a starting dose of 30 mg/day that could be reduced to 15 mg/day for tolerability and subsequently increased to 30 mg/day for clinical response.

Last observation carried forward (LOCF).

Y-MRS: Young Mania Rating Scale (range 0 to 60).

ABILIFY is indicated for the treatment of acute manic and mixed episodes associated with Bipolar I Disorder.

Please see IMPORTANT SAFETY INFORMATION, including **Bolded WARNING**, and INDICATIONS on page 4.



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. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular or infectious in nature.

ABILIFY is not approved for the treatment of patients with dementia-related psychosis.

Please see IMPORTANT SAFETY INFORMATION and INDICATIONS on page 4.





HELP ILLUMINATE THE PERSON WITHIN

IMPORTANT SAFETY INFORMATION and INDICATIONS for ABILIFY® (aripiprazole)

IMPORTANT SAFETY INFORMATION:

- 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. Although the causes of death were varied, most of the deaths appeared to be either cardiovascular or infectious in nature. ABILIFY (aripiprazole) is not approved for the treatment of patients with dementia-related psychosis (see Boxed WARNING).
- ABILIFY is contraindicated in patients with a known hypersensitivity to the product.
- As with all antipsychotic medications, including ABILIFY, a rare condition referred to as neuroleptic malignant syndrome (NMS) has been reported. As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of tardive dyskinesia (TD).
- Cerebrovascular adverse events (eg, stroke, transient ischemic attack), including fatalities, have been reported at an increased incidence in clinical trials of elderly patients with dementia-related psychosis treated with ABILIFY, including a significant dose-response relationship in a fixed-dose trial. ABILIFY is not approved for the treatment of patients with dementia-related psychosis.
- Hyperglycemia, including some serious cases ranging from ketoacidosis, hyperosmolar coma, or death, has been reported in patients treated with atypical antipsychotics. Patients on ABILIFY should be appropriately tested before and monitored during treatment.

ABILIFY may be associated with **orthostatic hypotension** and should be used with caution in patients with known cardiovascular disease, cerebrovascular disease, or conditions which would predispose them to hypotension.

As with other antipsychotic drugs, ABILIFY should be used with caution in patients with a history of seizures or with conditions that lower the seizure threshold.

Like other antipsychotics, ABILIFY may have the potential to impair judgment, thinking, or motor skills. Patients should not drive or operate hazardous machinery until they are certain ABILIFY does not affect them adversely.

Disruption of the body's ability to reduce core body temperature has been attributed to antipsychotics. Appropriate care is advised for patients who may exercise strenuously, be exposed to extreme heat, receive concomitant medication with anticholinergic activity, or be subject to dehydration.

As antipsychotics have been associated with esophageal dysmotility and aspiration, ABILIFY should be used cautiously in patients at risk for aspiration pneumonia.

As the possibility of a suicide attempt is inherent in psychotic illness and bipolar disorder, close supervision of high-risk patients should accompany drug therapy. Prescriptions for ABILIFY (aripiprazole) should be written for the smallest quantity consistent with good patient management to reduce the risk of overdose.

Physicians should determine if a patient is **pregnant** or intends to become pregnant while taking ABILIFY. Patients should be advised not to breast-feed while taking ABILIFY.

Patients should be advised to avoid alcohol while taking ABILIFY.

Both CYP3A4 and CYP2D6 are responsible for ABILIFY metabolism. Agents that induce CYP3A4 (eg, carbamazepine) could cause an increase in ABILIFY clearance and lower blood levels. Inhibitors of CYP3A4 (eg, ketoconazole) or CYP2D6 (eg, quinidine, fluoxetine, or paroxetine) can inhibit ABILIFY elimination and cause increased blood levels.

Commonly observed adverse events reported with ABILIFY in 3-week bipolar mania trials at a ≥5% incidence for ABILIFY and at a rate at least twice the rate of placebo include, respectively, akathisia (15% vs 4%), constipation (13% vs 6%), and accidental injury (6% vs 3%).

Treatment-emergent adverse events reported with ABILIFY in short-term trials at an incidence ≥10% and greater than placebo, respectively, include headache (31% vs 26%), agitation (25% vs 24%), anxiety (20% vs 17%), insomnia (20% vs 15%), nausea (16% vs 12%), dyspepsia (15% vs 13%), somnolence (12% vs 8%), akathisia (12% vs 5%), lightheadedness (11% vs 8%), vomiting (11% vs 6%), and constipation (11% vs 7%).

The adverse events reported in a 26-week, double-blind schizophrenia trial comparing ABILIFY and placebo were generally consistent with those reported in the short-term, placebo-controlled schizophrenia trials, except for a higher incidence of tremor: 9% for ABILIFY vs 1% for placebo.

INDICATIONS: ABILIFY is indicated for the treatment of:

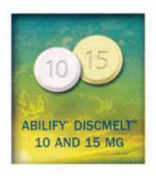
- Schizophrenia, including maintaining stability in patients who had been symptomatically stable on other antipsychotic medications for periods of 3 months or longer and observed for relapse during a period of up to 26 weeks*
- Acute manic and mixed episodes associated with Bipolar I Disorder
- Maintaining efficacy in patients with Bipolar I Disorder with a recent manic or mixed episode who had been stabilized and then maintained for at least 6 weeks*
- *Physicians who elect to use ABILIFY for extended periods should periodically re-evaluate the long-term usefulness of the drug for the individual patient.

ABILIFY is taken once daily with or without food.

Please see BRIEF SUMMARY of FULL PRESCRIBING INFORMATION, including **Boxed WARNING**, on the following pages.

New ABILIFY DISCMELT (aripiprazole)

Orally Disintegrating Tablets



Similar efficacy and safety to ABILIFY tablets

No liquid needed

Rapidly disintegrates

Convenient delivery of an effective dose



HELP ILLUMINATE THE PERSON WITHIN

ABILIFY® (aripiprazole) Tablets ABILIFY® DISCMELT™ (aripiprazole) **Orally Disintegrating Tablets ABILIFY®** (aripiprazole) Oral Solution

RONLY

Brief Summary of Prescribing Information. For complete prescribing information please consult official package circular.

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 (e.g., heart failure, sudden death) or infectious (e.g., peneumonia) in nature. ABILIFY (aripiprazole) is not approved for the treatment of patients with dementia-related psychosis.

CONTRAINDICATIONS

ABILIFY (aripiprazole) is contraindicated in patients with a known hypersensitivity to the product.

WARNINGS

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. ABILIFY is not approved for the treatment of patients with dementia-related psychosis (see Boxed WARNING).

Neuroleptic Malignant Syndrome (NMS)

Neuroleptic Malignant Syndrome (NMS) has A potentially fatal symptom complex sometimes referred to as Neuroleptic Malignant Syndrome (NMS) has been reported in association with administration of antipsychotic drugs, including aripiprazole. Two possible cases of NMS occurred during aripiprazole treatment in the premarketing worldwide clinical database. Clinical manifestations of NMS are hyperpyrexia, muscle rigidity, altered mental status, and evidence of autonomic instability (irregular pulse or blood pressure, tachycardia, diaphoresis, and control of the process of the

satus, and evidence or adultioning instability inequally pulse or blood pressure, tachycarda, diaphoresis, and cardiac dysrhythmia). Additional signs may include elevated creatine phosphokinase, myoglobinuria (rhabdomyolysis), and acute renal failure.

The diagnostic evaluation of patients with this syndrome is complicated. In arriving at a diagnosis, it is important to exclude cases where the clinical presentation includes both serious medical illness (e.g., pneumonia, systemic infection, etc) and untreated or inadequately treated extrapyramidal signs and symptoms (EPS). Other important considerations in the differential diagnosis include central anticholinergic twick that stocks drug fever and inference extraporates proving extraporate proving extraporate proving extraporates.

toxicity, heat stroke, drug fever, and primary central nervous system pathology.

The management of NMS should include: 1) immediate discontinuation of antipsychotic drugs and other drugs not essential to concurrent therapy; 2) intensive symptomatic treatment and medical monitoring; and 3) treatment of any concomitant serious medical problems for which specific treatments are available. There

is no general agreement about specific pharmacological treatment regimens for uncomplicated MMS.

If a patient requires antipsychotic drug treatment after recovery from NMS, the potential reintroduction of drug therapy should be carefully considered. The patient should be carefully monitored, since recurrences of NMS have been reported.

Tardive Dyskinesia

Tardive Dyskinesia
A syndrome of potentially irreversible, involuntary, dyskinetic movements may develop in patients treated with
antipsychotic drugs. Although the prevalence of the syndrome appears to be highest among the elderly,
especially elderly women, it is impossible to rely upon prevalence estimates to predict, at the inception of
antipsychotic treatment, which patients are likely to develop the syndrome. Whether antipsychotic drug
products differ in their potential to cause tardive dyskinesia is unknown.

The risk of developing tardive dyskinesia and the likelihood that it will become irreversible are believed to increase as the duration of treatment and the total cumulative dose of antipsychotic drugs administered to the patient increase. However, the syndrome can develop, although much less commonly, after relatively brief treatment periods at low doses.

There is no known treatment for established cases of tardive dyskinesia, although the syndrome may

remit, partially or completely, if antipsychotic treatment is withdrawn. Antipsychotic treatment, itself, however, may suppress (or partially suppress) the signs and symptoms of the syndrome and, thereby, may possibly mask the underlying process. The effect that symptomatic suppression has upon the long-term course of the syndrome is unknown.

course or the syndrome is unknown.

Given these considerations, ABILIPY should be prescribed in a manner that is most likely to minimize the occurrence of tardive dyskinesia. Chronic antipsychotic treatment should generally be reserved for patients who suffer from a chronic illness that (1) is known to respond to antipsychotic drugs, and (2) for whom alternative, equally effective, but potentially less harmful treatments are not available or appropriate. In patients who do require chronic treatment, the smallest dose and the shortest duration of treatment producing a satisfactory clinical response should be sought. The need for continued treatment should be reassessed periodically.

periodically.

If signs and symptoms of tardive dyskinesia appear in a patient on ABILIFY, drug discontinuation should be considered. However, some patients may require treatment with ABILIFY despite the presence of the syndrome.

Considered. However, some patients may require treatment with AbiLin't despite the presence of the syndrome. Cerebrovascular Adverse Events, including Stroke, in Elderly Patients with Dementia-Related Psychosis. In placebo-controlled clinical studies (two flexible dose and one fixed dose study) of dementia-related psychosis, there was an increased incidence of cerebrovascular adverse events (e.g., stroke, transient ischemic attack), including fatalities, in arripiprazole-treated patients (mean age: 84 years; range: 78-88 years). In the fixed-dose study, there was a statistically significant dose response relationship for cerebrovascular adverse events in patients treated with arripiprazole. Aripiprazole is not approved for the treatment of patients with dementia-related psychosis. (See also Boxed Warkling, Warklings: Increased Mortality in Elderly Patients with Dementia-Related Psychosis, and PRECAUTIONS: Use in Patients with Concomitant Illness: Safety Experience in Elderly Patients with Psychosis Associated with Alzheimer's Disease.)

Hyperglycemia and Diabetes Mellitus

Hyperglycenia, in some case extreme and associated with ketoacidosis or hyperosmolar coma or death, has been reported in patients treated with atypical antipsychotics. There have been few reports of hyperglycenia in patients treated with ABILIFY, Although fewer patients have been treated with ABILIFY, it is not known if this patients treated with ABILIFY. Although fewer patients have been treated with ABILIFY, it is not known if this more limited experience is the sole reason for the paucity of such reports. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes mellitus in patients with schizophrenia and the increasing incidence of diabetes mellitus in the general population. Given these confounders, the relationship between atypical mitrosychotic use and hyperglycemia-related adverse events is not completely understood. However, epidemiological studies which did not include ABILIFY suggest an increased risk of treatment-emergent hyperglycemia-related adverse events in patients treated with the atypical antipsychotics included in these studies. Because ABILIFY was not marketed at the time these studies were performed, it is not known if ABILIFY is associated with this increased risk. Precise risk estimates for hyperglycemia-related adverse events in patients treated with atypical antipsychotics are not available. antipsychotics are not available.

antipsychotics are not available.

Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics should be monitored regularly for worsening of glucose control. Patients with risk factors for diabetes mellitus (e.g., obesity, family history of diabetes) who are starting treatment with atypical antipsychotics should undergo fasting blood glucose testing at the beginning of treatment and periodically during treatment. Any patient treated with atypical antipsychotics should be monitored for symptoms of hyperglycemia including polydipsia, polyuria, polyphagia, and weakness. Patients who develop symptoms of hyperglycemia during treatment with atypical antipsychotics should undergo fasting blood glucose testing. In some cases, hyperglycemia has resolved when the atypical antipsychotic was discontinued; however, some patients required continuation of anti-diabetic treatment despite discontinuation of the suspect drug.

PRECAUTIONS

General

Orthostatic Hypotension

Aripiprazole may be associated with orthostatic hypotension, perhaps due to its α_1 -adrenergic receptor antagonism. The incidence of orthostatic hypotension-associated events from five short-term, placebo-

controlled trials in schizophrenia (n=926) on ABILIFY (aripiprazole) included: orthostatic hypotension (placebo 1%, aripiprazole 1.9%), orthostatic lightheadedness (placebo 1%, aripiprazole 0.9%), and syncope (placebo 1%, aripiprazole 0.6%). The incidence of orthostatic hypotension-associated events from short-term) placebo-controlled trials in bipolar mania (n=597) on ABILIFY included: orthostatic hypotension debedo 0%, aripiprazole 0.7%), orthostatic lightheadedness (placebo 0.5%, aripiprazole 0.5%), and syncope (placebo 0.9%, aripiprazole 0.5%). The incidence of a significant orthostatic change in blood pressure (defined as a decrease of at least 30 mmHg in systotic blood pressure when changing from a supine to standing position) for aripiprazole was not statistically different from placebo (in schizophrenia: 14% among aripiprazole-treated patients and 12% among placebo-treated patients and 12% among placebo-treated patients and 2% among placebo-treated patients an

placebo-treated patients).

Aripiprazole should be used with caution in patients with known cardiovascular disease (history of myocardial infarction or ischemic heart disease, heart failure or conduction abnormalities), cerebrovascular disease, or conditions which would predispose patients to hypotension (dehydration, hypovolemia, and treatment with antihypertensive medications).

Seizure
Seizure occurred in 0.1% (1/926) of aripiprazole-treated patients with schizophrenia in short-term, placebocontrolled trials. In short-term, placebo-controlled clinical trials of patients with bipolar mania, 0.3% (2/597)
of aripiprazole-treated patients and 0.2% (1/436) of placebo-treated patients experienced seizures. As with
other antipsychotic drugs, aripiprazole should be used cautiously in patients with a history of seizures or with
conditions that lower the seizure threshold, e.g., Alzheimer's dementia. Conditions that lower the seizure
threshold may be more prevalent in a population of 65 years or older.

Potential for Cognitive and Motor Impairment

Potential for Cognitive and Motor Impairment
In short-term, placebo-controlled trials of schizophrenia, somnolence was reported in 11% of patients on
BallLPY compared to 8% of patients on placebo; somnolence led to discontinuation in 0.1% (1/926) of
patients with schizophrenia on ABILIPY in short-term, placebo-controlled trials of bipolar mania, somnolence was reported in 14% of patients on ABILIPY compared to 7% optients with placebo, but did not lead to discontinuation of any patients with bipolar mania. Despite the
relatively modest increased incidence of somnolence compared to placebo, ABILIPY, like other antipsychotics,
may have the potential to impair judgment, thinking, or motor skills. Patients should be cautioned about
operating hazardous machinery, including automobiles, until they are reasonably certain that therapy with
ABILIPY does not affect them adversely.

Bary Temperature Benulation

Rody Temperature Regulation

Disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents.
Appropriate care is advised when prescribing aripiprazole for patients who will be experiencing conditions which may contribute to an elevation in core body temperature, e.g., exercising strenuously, exposure to extreme heat, receiving concomitant medication with anticholinergic activity, or being subject to dehydration.

Dyspriagia

Esophageal dysmotility and aspiration have been associated with antipsychotic drug use. Aspiration pneumonia is a common cause of morbidity and mortality in elderly patients, in particular those with advanced Alzheimer's dementia. Aripiprazole and other antipsychotic drugs should be used cautiously in patients at risk for aspiration pneumonia (see PRECAUTIONS: Use in Patients with Concomitant Illness).

The possibility of a suicide attempt is inherent in psychotic illnesses and bipolar disorder, and close supervision of high-risk patients should accompany drug therapy. Prescriptions for ABILIFY should be written for the smallest quantity consistent with good patient management in order to reduce the risk of overdose.

Use in Patients with Concomitant Illness
Clinical experience with ABILIFY in patients with certain concomitant systemic illnesses (see CLINICAL PHARMACOLOGY: Special Populations: Renal Impairment and Hepatic Impairment in Full Prescribing

ABILIFY has not been evaluated or used to any appreciable extent in patients with a recent history of myocardial infarction or unstable heart disease. Patients with these diagnoses were excluded from

myocardial infarction of unstable heart disease. Patients with these diagnoses were excluded from premarketing clinical studies.

Safety Experience in Elderly Patients with Psychosis Associated with Alzheimer's Disease: In three, 10-week, placebo-controlled studies of aripiprazole in elderly patients with psychosis associated with Alzheimer's disease (n=938; mean age: 82.4 years; range: 56-99 years), the treatment-emergent adverse events that were reported at an incidence of ≥3% and aripiprazole incidence at least twice that for placebo were asthenia (placebo 3%, aripiprazole 8%), somnolence (placebo 3%, aripiprazole 9%), uninterported dispersion (placebo 4%, aripiprazole 8%), somnolence (placebo 3%, aripiprazole 9%), uninterported dispersion (placebo 4%, aripiprazole 9%), uniterported dispersion (placebo 4%, aripi

were asthenia (placebo 3%, aripiprazole 8%), somnolence (placebo 3%, aripiprazole 9%), urinary incontinence (placebo 1%, aripiprazole 5%), excessive salivation (placebo 0%, aripiprazole 4%), and lightheadedness (placebo 1%, aripiprazole 4%).

The safety and efficacy of ABILIFY in the treatment of patients with psychosis associated with dementia have not been established. If the prescriber elects to treat such patients with ABILIFY, vigitance should be exercised, particularly for the emergence of difficulty swallowing or excessive somnolence, which could predispose to accidental injury or aspiration. (See also Boxed WARNING, WARNINGS: Increased Mortality in Elderty Patients with Dementia-Related Psychosis, and Cerebrovascular Adverse Events, Including Stroke, in Elderty Patients with Dementia-Related Psychosis.)

Information for Patients

Physicians are advised to discuss the following issues with patients for whom they prescribe ABILIFY:

Interference with Cognitive and Motor Performance: Because aripiprazole may have the potential to impair judgment, thinking, or motor skills, patients should be cautioned about operating hazardous machinery, including automobiles, until they are reasonably certain that aripiprazole therapy does not affect them adversely.

Pregnancy: Patients should be advised to notify their physician if they become pregnant or intend to become pregnant during therapy with ABILIFY.

Nursing: Patients should be advised not to breast-feed an infant if they are taking ABILIFY.

Concomitant Medication: Patients should be advised to inform their physicians if they are taking, or plan to take, any prescription or over-the-counter drugs, since there is a potential for interactions. Alcohol: Patients should be advised to avoid alcohol while taking ABILIFY.

Heat Exposure and Dehydration: Patients should be advised regarding appropriate care in avoiding overheating

Sugar Content: Patients should be advised that each mL of ABILIFY oral solution contains 400 mg of sucrose and 200 mg of fructose.

Phenylketonurics: Phenylalanine is a component of aspartame. Each ABILIFY DISCMELT orally disintegrating tablet contains the following amounts: 10 mg - 1.12 mg phenylalanine and 15 mg - 1.68 mg phenylalanine.

Drug-Drug Interactions

Given the primary CNS effects of aripiprazole, caution should be used when ABILIFY is taken in combination with other centrally acting drugs and alcohol. Due to its α_1 -adrenergic receptor antagonism, aripiprazole has the potential to enhance the effect of certain antihypertensive agents.

Potential for Other Drugs to Affect ABILIFY

the potential to enhance the effect of certain antihypertensive agents.

Potential for Other Drugs to Affect ABILIFY
Aripiprazole is not a substrate of CYP1A1, CYP1A2, CYP2A6, CYP2B6, CYP2C8, CYP2C9, CYP2C19, or CYP2E1
enzymes. Aripiprazole also does not undergo direct glucuronidation. This suggests that an interaction of
aripiprazole with inhibitors or inducers of these enzymes, or other factors, like smoking, is unlikely,
Both CYP3A4 and CYP2D6 are responsible for aripiprazole metabolism. Agents that induce CYP3A4 (e.g.,
carbamazepine) could cause an increase in aripiprazole clearance and lower blood levels. Inhibitors of CYP3A4 (e.g., ketoconazole) or CYP2D6 (e.g., quinding, fluoxetine, or paroxetine) can inhibit aripiprazole
elimination and cause increased blood levels.

Ketoconazole: Oadministration of ketoconazole (200 mg/day for 14 days) with a 15-mg single dose of
aripiprazole increased the AUC of aripiprazole and its active metabolite by 63% and 77%, respectively. The
effect of a higher ketoconazole dose (400 mg/day) has not been studied. When concomitant administration of ketoconazole with aripiprazole oscurs, aripiprazole dose should be reduced to one-half of its normal dose.
Other strong inhibitors of CYP3A4 (firaconazole) would be expected to have similar effects and need similar
dose reductions; weaker inhibitors of CYP3A6 (iraconazole) would be expected to have similar effects and need similar
dose reductions; weaker inhibitors of CYP2D6, increased the AUC of aripiprazole with quinidine (166 mg/day) for 13
days), a potent inhibitor of CYP2D6, increased the AUC of aripiprazole by 112% but decreased the AUC of its
active metabolite, dehydro-aripiprazole, by 35%. Aripiprazole dose should be reduced to one-half of its
normal dose when concomitant administration of quinidine with aripiprazole occurs. Other significant
inhibitors of CYP2D6, such as fluoxetine or paroxetine, would be expected to have similar effects and,
therefore, should be accompanied by similar dose reductions. When the CYP2D6 inhibi

Carbamazepine: Coadministration of carbamazepine (200 mg BID), a potent CYP3A4 inducer, with aripiprazole (30 mg QD) resulted in an approximate 70% decrease in $C_{\rm max}$ and AUC values of both aripiprazole and its active metabolite, dehydro-aripiprazole. When carbamazepine is added to aripiprazole therapy, aripiprazole dose should be doubled. Additional dose increases should be based on clinical evaluation. When carbamazepine is withdrawn from the combination therapy, aripiprazole dose should then

No clinically significant effect of famotidine, valproate, or lithium was seen on the pharmacokinetics of aripiprazole (see CLINICAL PHARMACOLOGY: Drug-Drug interactions in Full Prescribing Information).

Potential for ABILIFY (aripiprazole) to Affect Other Drugs

Aripiprazole is unlikely to cause clinically important pharmacokinetic interactions with drugs metabolized by cytochrome P450 enzymes. In *in vivo* studies, 10- to 30-mg/day doses of aripiprazole had no significant effect on metabolism by CYP2D6 (dextromethorphan), CYP2C9 (warfarin), CYP2C9 (merazole, warfarin), and CYP3A4 (dextromethorphan) substrates. Additionally, aripiprazole and dehydro-aripiprazole did not show potential for altering CYP1A2-mediated metabolism *in vitro* (see CLINICAL PHARMACOLOGY: Drug-Drug Interactions in Full Prescribing Information).

Alcohol: There was no significant difference between aripiprazole coadministered with ethanol and placebo coadministered with ethanol on performance of gross motor skills or stimulus response in healthy subjects. As with most psychoactive medications, patients should be advised to avoid alcohol while taking ABILIFY.

Carcinogenesis, Mutagenesis, Impairment of Fertility

Cardingeresis. Lifetime cardinogenicity studies were conducted in ICR mice and in Sprague-Dawley (SD) and F344 rats. Aripiprazole was administered for 2 years in the diet at doses of 1, 3, 10, and 30 mg/kg/day to ICR mice and 1, 3, and 10 mg/kg/day to F344 rats (0.2 to 5 and 0.3 to 3 times the maximum recommended human dose 1, 3, and 10 mg/kg/day to F344 rats (0.2 to 5 and 0.3 to 3 times the maximum recommended human dose MRHD] based on mg/m², respectively). In addition, SD rats were dosed orally for 2 years at 10, 20, 40, and 60 mg/kg/day (3 to 19 times the MRHD based on mg/m²). Aripiprazole did not induce tumors in male mice or rats. In female mice, the incidences of pituitary gland adenomas and mammary gland adenocarcinomas and denocarcinomas and adenocarcinomas are denocarcinomas and sensor at MRHD based on AUC and 0.5 to 5 times the MRHD based on mg/m²). In female rats, the incidence of mammary gland fibroadenomas was increased at a dietary dose of 10 mg/kg/day (0.1 times human exposure at MRHD based on AUC and 3 times the MRHD based on mg/m²); and the incidences of addrenocortical carcinomas and combined adrenocortical adenomas/carcinomas were increased at an oral dose of 60 mg/kg/day (14 times human exposure at MRHD based on AUC and 19 times the MRHD based on mg/m²). Proliferative changes in the nitivitary and mammary gland of rincients have been observed following chronic process.

Proliferative changes in the pituitary and mammary gland of rodents have been observed following chronic administration of other antipsychotic agents and are considered prolactin-mediated. Serum prolactin was not measured in the aripiprazole carcinogenicity studies. However, increases in serum prolactin levels were observed in female mice in a 13-week dietary study at the doses associated with mammary gland and pituitary tumors. Serum prolactin was not increased in female rats in 4- and 13-week dietary studies at the dose associated with mammary gland tumors. The relevance for human risk of the findings of prolactin-mediated endocrine tumors in ordents in unpravail. endocrine tumors in rodents is unknown.

Mutagenesis

Mutagenesis
The mutagenic potential of aripiprazole was tested in the *in vitro* bacterial reverse-mutation assay, the *in vitro* bacterial DNA repair assay, the *in vitro* forward gene mutation assay in mouse lymphoma cells, the *in vitro* chromosomal aberration assay in chinese hamster lung (CHL) cells, the *in vivo* micronucleus assay in mitor, and the unscheduled DNA synthesis assay in rats. Aripiprazole and a metabolite (2.3-DCPP) were clastogenic in the *in vitro* chromosomal aberration assay in CHL cells with and without metabolic activation. The metabolite, 2.3-DCPP, produced increases in numerical aberrations in the *in vitro* assay in CHL cells in the absence of metabolic activation. A positive response was obtained in the *in vivo* micronucleus assay in mice; however, the response was shown to be due to a mechanism not considered relevant to humans.

Impairment or returned with oral doses of 2, 6, and 20 mg/kg/day (0.6, 2, and 6 times the maximum recommended human dose [MRHD] on a mg/m² basis) of aripiprazole from 2 weeks prior to mating through day 7 of gestation. Estrus cycle irregularities and increased corpora lutea were seen at all doses, but no impairment of fertility was seen. Increased pre-implantation loss was seen at 6 and 20 mg/kg, and decreased

Impairment or returnly was seen, incleased pre-impairment or so was seen at 0 and 20 mg/kg, and decleased fetal weight was seen at 20 mg/kg.

Male rats were treated with oral doses of 20, 40, and 60 mg/kg/day (6, 13, and 19 times the MRHD on a mg/m² basis) of aripinzacie from 9 weeks prior to mating through mating, Disturbances in spermatogenesis were seen at 60 mg/kg, and prostate atrophy was seen at 40 and 60 mg/kg, but no impairment of fertility

Pregnancy

Pregnancy Category C

In animal studies, aripiprazole demonstrated developmental toxicity, including possible teratogenic effects in

Pregnant rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the maximum recommended human dose [MRHID] on a mg/m² basis) of aripiprazole during the period of organogenesis Gestation was slightly prolonged at 30 mg/kg. Treatment caused a slight delay in fetal development, as evidenced by decreased fetal weight (30 mg/kg), undescended testes (30 mg/kg), and delayed skeletal ossification (10 and 30 mg/kg). There were no adverse effects on embryofetal or pup survival. Delivered offspring had decreased bodyweights (10 and 30 mg/kg), and increased incidences of hepatodiaphragmatic nodules and diaphragmatic hernia at 30 mg/kg (the other dose groups were not examined for these findings). A low incidence of diaphragmatic hernia was also seen in the fetuses exposed to 30 mg/kg). Postnatally, delayed vaginal opening was seen at 10 and 30 mg/kg and impaired reproductive performance (decreased fertility rate, corpora lutea, implants, and live fetuses, and increased post-implantation loss, likely mediated through effects on female offspring) was seen at 30 mg/kg. Some maternal toxicity was seen at 30 mg/kg, however, there was no evidence to suggest that these developmental effects were secondary to maternal toxicity. Pregnant rabbits were treated with oral doses of 10, 30, and 100 mg/kg/dy, 2, 3, and 11 mse human exposure at MRHD based on AUC and 6, 19, and 65 times the MRHD based on mg/m²) of aripiprazole during the period of organogenesis. Decreased maternal food consumption and increased abortions were seen at 100 mg/kg), increased incidence of skeletal abnormality (fused sternebrae at 30 and 100 mg/kg) and minor skeletal variations (100 mg/kg), and minor skeletal variations (100 mg/kg). Pregnant rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the maximum

mg/kg), increased incidence of skeletal abnormality (fused sternebrae at 30 and 100 mg/kg) and minor skeletal variations (100 mg/kg).

In a study in which rats were treated with oral doses of 3, 10, and 30 mg/kg/day (1, 3, and 10 times the MRHD on a mg/m² basis) of aripiprazole perinatally and postnatally (from day 17 of gestation through day 21 postpartum), slight maternal toxicity and slightly prolonged gestation were seen at 30 mg/kg. An increase in stillibitries, and decreases in pup weight (persisting into adulthoot) and survival, were seen at this dose. There are no adequate and well-controlled studies in pregnant women. It is not known whether aripiprazole can cause fetal harm when administered to a pregnant woman or can affect reproductive capacity. Aripiprazole should be used during pregnancy only if the potential benefit outweighs the potential risk to the fetus.

Labor and Delivery

The effect of aripiprazole on labor and delivery in humans is unknown

Nursing Mothers

Aripiprazole was excreted in milk of rats during lactation. It is not known whether aripiprazole or its metabolites are excreted in human milk. It is recommended that women receiving aripiprazole should not breast-feed.

Pediatric Use

Safety and effectiveness in pediatric and adolescent patients have not been established.

Geriatric Use

Of the 7951 patients treated with aripiprazole in premarketing clinical trials, 991 (12%) were ≥65 years old
and 789 (10%) were ≥75 years old. The majority (88%) of the 991 patients were diagnosed with dementia of

and 798 (10%) were 275 years out. The majority (60%) of the 991 patients were diagnosed with definition of the Alzheimer's type.

Placebo-controlled studies of aripiprazole in schizophrenia or bipolar mania did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. There was no effect of age on the pharmacokinetics of a single 15-mg dose of aripiprazole. Aripiprazole clearance was decreased by 20% in elderly subjects (≥65 years) compared to younger adult subjects (18 to 64 years), but there was no detectable effect of age in the population pharmacokinetic analysis in exhibitopheria antients.

b4 years), but there was no detectable effect of age in the population plantacounted sharped schizophrenia patients.

Studies of elderly patients with psychosis associated with Alzheimer's disease have suggested that the may be a different tolerability profile in this population compared to younger patients with schizophrenia (see Boxed WARNING; WARNINGS: Increased Mortality in Elderly Patients with Dementia-Related Psychosis; Cerebrovascular Adverse Events, Including Stroke, in Elderly Patients with Dementia-

Related Psychosis: and PRECAUTIONS: Use in Patients with Concomitant Illness). The safety and efficacy of ABILIFY (aripiprazole) in the treatment of patients with psychosis associated with Alzheimer's disease has been established. If the prescriber elects to treat such patients with ABILIFY, vigilance should be exercised

ADVERSE REACTIONS

Aripiprazole has been evaluated for safety in 7951 patients who participated in multiple-dose, premarketing trials in schizophrenia, bipolar mania, and dementia of the Alzheimer's type, and who had approximately 5235

trials in schizophrenia, bipolar mania, and dementia of the Alzheimer's type, and who had approximately 5235 patient-years of exposure. A total of 2280 aripiparacle-treated patients were treated for at least 180 days and 1558 aripiprazole-treated patients had at least 1 year of exposure. The conditions and duration of treatment with aripiprazole included (in overlapping categories) double-blind, comparative and noncomparative open-label studies, inpatient and outpatient studies, fixed-and flexible-dose studies, and short- and longer-term exposure. Adverse events during exposure were obtained by collecting voiunteered adverse events, as well as results of physical examinations, vital signs, weights, laboratory analyses, and ECG. Adverse experiences were recorded by cinical investigators using terminology of their own choosing. In the tables and tabulations that follow, modified COSTART dictionary terminology has been used initially to classify reported adverse events into a smaller number of standardized event categories, in order to provide a meaningful estimate of the proportion of individuales experiencing adverse events.

events into a smaller manner or standardized event categories, in order to provide a meaningful estimate or the proportion of individuals experiencing adverse events.

The stated frequencies of adverse events represent the proportion of individuals who experienced at least once, a treatment-emergent adverse event of the type listed. An event was considered treatment emergent if occurred for the first time or worsened while receiving therapy following baseline evaluation. There was no attempt to use investigator causality assessments; ie, all reported events are included.

The prescriber should be aware that the figures in the tables and tabulations cannot be used to predict the included.

The prescriber should be aware that the lightes in the tables and accutations cannot be used to predict the incidence of side effects in the course of usual medical practice where patient characteristics and other factors differ from those that prevailed in the clinical trials. Similarly, the cited frequencies cannot be compared with figures obtained from other clinical investigations involving different treatment, uses, and investigators. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and nondrug factors to the adverse event incidence in the population studied.

Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Schizophrenia The following findings are based on a pool of five placebo-controlled trials (four 4-week and one 6-week) in which aripiprazole was administered in doses ranging from 2 to 30 mg/day.

Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials Overall, there was no difference in the incidence of discontinuation due to adverse events between aripiprazole-treated (7%) and placebo-treated (9%) patients. The types of adverse events that led to discontinuation were similar between the aripiprazole- and placebo-treated patients.

Adverse Findings Observed in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Mania The following findings are based on a pool of 3-week, placebo-controlled, bipolar mania trials in which aripiprazole was administered at doses of 15 or 30 mg/day.

Adverse Events Associated with Discontinuation of Treatment in Short-Term, Placebo-Controlled Trials
Overall, in patients with bipolar mania, there was no difference in the incidence of discontinuation due to
adverse events between aripiprazole-treated (11%) and placebo-treated (9%) patients. The types of adverse
events that led to discontinuation were similar between the aripiprazole and placebo-treated patients.

Commonly Observed Adverse Events in Short-Term, Placebo-Controlled Trials of Patients with Bipolar Mania Commonly observed adverse events associated with the use of aripiprazole in patients with bipolar mania (incidence of 5% or greater and aripiprazole incidence at least twice that for placebo) are shown in Table 1. There were no adverse events in the short-term trials of schizophrenia that met these criteria.

Table 1: Commonly Observed Adverse Events in Short-Term,
Placebo-Controlled Trials of Patients with Bipolar Mania

	Percentage of Patier	nts Reporting Event	
Adverse Event	Aripiprazole (n=597)	Placebo (n=436)	
Accidental Injury	6	3	
Constipation	13	6	
Akathisia	15	4	

Adverse Events Occurring at an Incidence of 2% or More Among Aripiprazole-Treated Patients and Greater than Placebo in Short-Term, Placebo-Controlled Trials

Table 2 enumerates the pooled incidence, rounded to the nearest percent, of treatment-emergent adverse events that occurred during acute therapy (up to 6 weeks in schizophrenia and up to 3 weeks in bipolar mania), including only those events that occurred in 2% or more of patients treated with aripiprazole (doses $\ge 2 \operatorname{my/day}$) and for which the incidence in patients treated with packed with packed in the combined dataset. patients treated with placebo in the combined dataset.

Table 2: Treatment-Emergent Adverse Events in Short-Term Placeho-Controlled Trials

	Percentage of Patien	ts Reporting Event ^a	
Body System Adverse Event	Aripiprazole (n=1523)	Placebo (n=849)	
Body as a Whole			
Headache	31	26	
Asthenia	8	7	
Accidental Injury	5	4	
Peripheral Edema	2	1	
Cardiovascular System			
Hypertension	2	1	
Digestive System			
Nausea	16	12	
Dyspepsia	15	13	
Vomiting	11	6	
Constipation	11	7	
Musculoskeletal System			
Myalqia	4	3	
Nervous System	•	ŭ	
Agitation	25	24	
Anxiety	20	17	
Insomnia	20	15	
Somnolence	12	8	
Akathisia	12	5	
Lightheadedness	11	8	
Extrapyramidal Syndrome	6	4	
Tremor	4	3	
Increased Salivation	3	1	
Respiratory System			
Pharyngitis	4	3	
Rhinitis	4	3 2	
Coughing	3	2	
Special Senses			
Blurred Vision	3	1	

^a Events reported by at least 2% of patients treated with aripiprazole, except the following events, which had an incidence equal to or less than placebo: abdominal pain, back pain, dental pain, diarrhea, dry mouth, anorexia, psychosis, hypertonia, upper respiratory tract infection, rash, vaglinitis¹, dysmenorrhea⁴.

f Percentage based on gender total.

An examination of population subgroups did not reveal any clear evidence of differential adverse event incidence on the basis of age, gender, or race.

Dose-Related Adverse Events

Schizophrenia

Schizophrenia Dose response relationships for the incidence of treatment-emergent adverse events were evaluated from four trials in patients with schizophrenia comparing various fixed doses (2, 10, 15, 20, and 30 mg/day) of aribiprazole to placebo. This analysis, stratified by study, indicated that the only adverse event to have a possible dose response relationship, and then most prominent only with 30 mg, was somnolence (placebo, 7.7%; 15 mg, 8.7%; 20 mg, 7.5%; 30 mg, 15.3%).

Extrapyramidal Symptoms

Extrapyramidal Symptoms
In the short-term, placebo-controlled trials of schizophrenia, the incidence of reported EPS for aripiprazole-treated patients was 6% vs. 6% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of reported EPS-related events excluding events related to akathisia for aripiprazole-treated patients was 17% vs. 12% for placebo. In the short-term, placebo-controlled trials in bipolar mania, the incidence of akathisia-related events for aripiprazole-treated patients was 15% vs. 4% for placebo. Objectively collected data from those trials was collected on the Simpson Angus Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia) and the Assessments of involuntary Movement Scales (for Schiknesias). In the schizophrenia trials, the objectively collected data did not show a difference between aripiprazole and placebo, with the exception of the Barnes Akathisia Scale (aripiprazole, 0.08; placebo, -0.05). In the bipolar mania trials, the Simpson Angus Rating Scale and the Barnes Akathisia Scale showed a significant difference between aripiprazole and placebo (aripiprazole, 0.61; placebo, 0.03 and aripiprazole, 0.25; placebo, -0.06). Changes in the Assessments of Involuntary Movement Scales were similar for the aripiprazole and placebo groups.

Similarly, in a long-term (26-week), placebo-controlled trial of schizophrenia, objectively collected data on the Simpson Angus Rating Scale (for EPS), the Barnes Akathisia Scale (for akathisia), and the Assessments of Involuntary Movement Scales (for dyskinesias) did not show a difference between aripiprazole and placebo.

Laboratory Test Abnormalities

Laboratory lest Andromanuse.

A between group comparison for 3- to 6-week, placebo-controlled trials revealed no medically important differences between the arripiprazole and placebo groups in the proportions of patients experiencing potentially clinically significant changes in routine serum chemistry, hematology, or urinalysis parameters. Similarly, there were no arripiprazole/placebo differences in the incidence of discontinuations for changes in serum chemistry,

were no anyprazous/praced unretences in the incidence of discontinuations for changes in serum chemistry, hematology, or urinalysis.

In a long-term (26-week), placebo-controlled trial there were no medically important differences between the aripiprazoie and placebo patients in the mean change from baseline in prolactin, fasting glucose, triglyceride, HDL, LDL, and total cholesterol measurements.

Weight Gain

Weight Gain
In 4- to 6-week trials in schizophrenia, there was a slight difference in mean weight gain between
aripiprazole and placebo patients (+0.7 kg vs. -0.05 kg, respectively), and also a difference in the proportion
of patients meeting a weight gain criterion of ≥7% of body weight [aripiprazole (8%) compared to placebo
(3%)]. In 3-week trials in mania, the mean weight gain for aripiprazole and placebo patients was 0.0 kg vs.
-0.2 kg, respectively. The proportion of patients meeting a weight gain criterion of ≥7% of body weight was
aripiprazole (3%) compared to placebo (2%).
Table 3 provides the weight change results from a long-term (26-week), placebo-controlled study of
aripiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of
≥7% of body weight relative to baseline, categorized by BMI at baseline:

Table 3: Weight Change Results Categorized by BMI at Baseline: Placebo-Controlled Study in Schizophrenia, Safety Sam

	Bl	VII <23	BM	11 23-27	BMI	>27
	Placebo	Aripiprazole	Placebo	Arlpiprazole	Placebo	Aripiprazole
Mean change from baseline (kg)	-0.5	-0.5	-0.6	-1.3	-1.5	-2.1
% with ≥7% increase BW	3.7%	6.8%	4.2%	5.1%	4.1%	5.7%

Table 4 provides the weight change results from a long-term (52-week) study of aripiprazole, both mean change from baseline and proportions of patients meeting a weight gain criterion of \geq 7% of body weight relative to baseline, categorized by BMI at baseline:

Table 4: Weight Change Results Categorized by BMI at Baseline: Active-Controlled Study in Schizophrenia, Safety Sample

	BMI <23	BMI 23-27	BMI >27
Mean change from baseline (kg)	2.6	1.4	-1.2
% with ≥7% increase BW	30%	19%	8%

Between group comparisons for a pooled analysis of placebo-controlled trials in patients with schizophrenia or bipolar mania, revealed no significant differences between aripiprazole and placebo in the proportion of patients experiencing potentially important changes in ECG parameters. Aripiprazole was associated with a median increase in heart rate of 5 beats per minute compared to a 1 beat per minute increase among placebo patients.

Additional Findings Observed in Clinical Trials

Adverse Events in Long-Term, Double-Blind, Placebo-Controlled Trials

The adverse events reported in a 26-week, double-blind trial comparing ABILIFY (aripiprazole) and placebo in patients with schizophrenia were generally consistent with those reported in the short-term, placebo-controlled trials, except for a higher incidence of tremor [9% (13/153) for ABILIFY vs. 1% (2/153) for placebo]. In this study, the majority of the cases of tremor were of mild intensity (9/13 mild and 4/13 moderate), occurred early in therapy (9/13 <4/d>
49(3), and were of limited duration (9/13 <10 days). Tremor inferouently led to discontinuation (<1%) of ABILIFY. In addition, in a long-term (52-week), active-controlled study, the incidence of tremor for ABILIFY was 4% (34/859). A similar adverse event profile was observed in a long-term study in binalor disorder. study in bipolar disorder.

study in bipolar disorder.

Other Adverse Events Observed During the Premarketing Evaluation of Aripiprazole
Following is a list of modified COSTART terms that reflect treatment-emergent adverse events as defined in the introduction to the ADVERSE REACTIONS section reported by patients treated with aripiprazole at multiple doses ≥2 mg/day during any phase of a trial within the database of 7951 patients. All reported events are included except those already listed in Table 2, or other parts of the ADVERSE REACTIONS section, those considered in the WARNINGS or PRECAUTIONS, those event terms which were so general as to be uninformative, events reported with an incidence of ≈0.05% and which did not have a substantly of being acutely life-threatening, events that are otherwise common as background events, and events considered unlikely to be drug related. It is important to emphasize that, although the events reported occurred during treatment with aripiprazole, they were not necessarily caused by it.

Events are further categorized by body system and listed in order of decreasing frequency according to the following definitions: frequent adverse events are those occurring in at least 1/100 patients (only those not already listed in the tabulated results from placebo-controlled trials appear in this listing); infrequent adverse events are those occurring in fewer than 1/1000 patients.

1/1000 patients.

Body as a Whole: Frequent – flu syndrome, fever, chest pain, rigidity (including neck and extremity), neck pain, pelvic pain; Intrequent – face edema, suicide attempt, malaise, migraine, chills, photosensitivity, tightness (including abdomen, back, extremity, head, jaw, neck, and tongue), jaw pain, bloatien, enlarged abdomen, chest tightness, throat pain; Rare – moniliasis, head heaviness, throat tightness, Mendelson's syndrome, heat stroke.

Cardiovascular System: Frequent – tachycardia (Including ventricular and supraventricular), hypotension.

Cardiovascular System: Frequent — tachycardia (including ventricular and supraventricular), hypotension radycardia; Infrequent — pabiptation, hemorrhage, heart failure, myocardial infarction, cardiac rest, atrial fibrillation, AV block, prolonged QT interval, extrasystoles, myocardial ischemia, deep vein thrombosis, angina pectoris, pallor, cardiopulmonary arrest, phlebitis; Rare — bundle branch block, atrial flutter, vasovagal reaction, cardiomegaly, thrombophlebitis, cardiopulmonary failure.

Digestive System: Frequent — nausea and vomitting; Infrequent — increased appetite, dysphagia, gastroenteritis, flatulence, toott caries, gastritis, ginglivitis, gastronistinal hemorrhage, shomatris, collitis, tongue edema, cholecystitis, mouth ulcer, oral moniliasis, eructation, fecal impaction, cholelithiasis; Rare —

esophagitis, hematemesis, intestinal obstruction, gum hemorrhage, hepatitis, peptic ulcer, glossitis, melena,

duodenal ulcer, cheilitis, hepatomegaly, pancreatitis.

Endocrine System: Infrequent -- hypothyroidism; Rare -- goiter, hyperthyroidism.

Hemic/Lymphatic System: Frequent - ecchymnosis, anemia; Infrequent - hypochromic anemia, leukocytosis, leukopenia (including neutropenia), lymphadenopathy, eosinophilia, macrocytic anemia; Rare thrombocythemia, thrombocytopenia, petechiae.

Metabolic and Nutritional Disorders: Frequent - weight loss, creating phosphokinase increased. dehydration; Infrequent – edema, hyperglycemia, hypercholesteremia, hypokalemia, diabetes mellitus, hypoglycemia, hyperlipemia, SGPT increased, thirst, BUN increased, hyponatremia, SGOT increased, creatinine increased, cyanosis, alkaline phosphatase increased, bilirubinemia, iron deficiency anemia, hyperkalemia, hyperuricemia, obesity; Rare - lactic dehydrogenase increased, hypernatremia, gout, hypoglycemic reaction.

Musculoskeletal System: Frequent - muscle cramp; Infrequent - arthralgia, myasthenia, arthrosis, bone pain, arthritis, muscle weakness, spasm, bursitis, myopathy; Rare - rheumatoid arthritis, rhabdomyolysis, tendonitis, tenosynovitis.

Nervous System: Frequent – depression, nervousness, schizophrenic reaction, hallucination, hostility, confusion, paranoid reaction, suicidal thought, abnormal gait, manic reaction, delusions, abnormal dream; Infrequent - emotional lability, twitch, cogwheel rigidity, impaired concentration, dystonia, vasodilation, paresthesia, impotence, extremity tremor, hypesthesia, vertigo, stupor, bradykinesia, apathy, panic attack, decreased libido, hypersomnia, dyskinesia, manic depressive reaction, ataxia, visual hallucination, cerebrovascular accident, hypokinesia, depersonalization, impaired memory, delirium, dysarthria, tardive dyskinesia, amnesia, hyperactivity, increased libido, myoclonus, restless leg, neuropathy, dysphoria, hyperkinesia, cerebral ischemia, increased reflexes, akinesia, decreased consciousness, hyperesthesia, slowed thinking; Rare – blunted affect, euphoria, incoordination, oculogyric crisis, obsessive thought, hypotonia, buccoglossal syndrome, decreased reflexes, derealization, intracranial hemorrhage.

*Respiratory System: Frequent – sinusitis, dyspnea, pneumonia, asthma; Infrequent – epistaxis, hiccup,

laryngitis, aspiration pneumonia; Rare - pulmonary edema, increased sputum, pulmonary embolism, hypoxia,

respiratory failure, apnea, dry nasal passages, hemoptysis.

Skin and Appendages: Frequent – skin ulcer, sweating, dry skin; Infrequent – pruritus, vesiculobullous rash, acne, eczema, skin discoloration, alopecia, seborrhea, psoriasis; Rare - maculopapular rash, exfoliative dermatitis, urticaria

Special Senses: Frequent - conjunctivitis; Infrequent - ear pain, dry eye, eye pain, tinnitus, cataract, otitis media, altered taste, blepharitis, eye hemorrhage, deafness; Rare - diplopia, frequent blinking, ptosis, otitis externa, amblyopia, photophobia.

Urogenital System: Frequent – urinary incontinence; Infrequent – urinary frequency, leukorrhea, urinary retention, cystitis, hematuria, dysuria, amenorrhea, vaginal hemorrhage, abnormal ejaculation, kidney failure, vaginal moniliasis, urinary urgency, gynecomastia, kidney calculus, albuminuria, breast pain, urinary burning; Rare – nocturia, polyuria, menorrhagia, anorgasmy, glycosuria, cervicitis, uterus hemorrhage, female lactation, urolithiasis, priapism.

Other Events Observed During the Postmarketing Evaluation of Aripiprazole

Voluntary reports of adverse events in patients taking aripiprazole that have been received since market introduction and not listed above that may have no causal relationship with the drug include rare occurrences of allergic reaction (e.g., anaphylactic reaction, angioedema, laryngospasm, pruritus, or urticaria)

DRUG ABUSE AND DEPENDENCE

Controlled Substance

ABILIFY (aripiprazole) is not a controlled substance.

Abuse and Dependence

Aripiprazole has not been systematically studied in humans for its potential for abuse, tolerance, or physical dependence. In physical dependence studies in monkeys, withdrawal symptoms were observed upon abrupt cessation of dosing. While the clinical trials did not reveal any tendency for any drug-seeking behavior, these observations were not systematic and it is not possible to predict on the basis of this limited experience the extent to which a CNS-active drug will be misused, diverted, and/or abused once marketed. Consequently, patients should be evaluated carefully for a history of drug abuse, and such patients should be observed closely for signs of ABILIFY misuse or abuse (e.g., development of tolerance, increases in dose, drug-seeking behavior).

OVERDOSAGE

MedDRA terminology has been used to classify the adverse events.

Human Experience

A total of 76 cases of deliberate or accidental overdosage with aripiprazole have been reported worldwide. These include overdoses with aripiprazole alone and in combination with other substances. No fatality was reported from these cases. Of the 44 cases with known outcome, 33 recovered without sequelae and one recovered with sequelae (mydriasis and feeling abnormal). The largest known acute ingestion with a known outcome involved 1080 mg of aripiprazole (36 times the maximum recommended daily dose) in a patient who fully recovered, included in the 76 cases are 10 cases of deliberate or accidental overdosage in children (age 12 and younger) involving aripiprazole ingestions up to 195 mg with no fatalities.

Common adverse events (reported in at least 5% of all overdose cases) reported with aripiprazole overdosage (alone or in combination with other substances) include vomiting, somnolence, and tremor. Other clinically important signs and symptoms observed in one or more patients with aripiprazole overdoses (alone or with other substances) include acidosis, aggression, aspartate aminotransferase increased, atrial fibrillation, bradycardia, coma, confusional state, convulsion, blood creatine phosphokinase increased, depressed level of consciousness, hypertension, hypokalemia, hypotension, lethargy, loss of consciousness, QRS complex prolonged, QT prolonged, pneumonia aspiration, respiratory arrest, status epilepticus, and tachycardia.

Management of Overdosage

No specific information is available on the treatment of overdose with aripiprazole. An electrocardiogram should be obtained in case of overdosage and, if QTc interval prolongation is present, cardiac monitoring should be instituted. Otherwise, management of overdose should concentrate on supportive therapy maintaining an adequate airway, oxygenation and ventilation, and management of symptoms. Close medical supervision and monitoring should continue until the patient recovers.

Charcoal: In the event of an overdose of ABILIFY, an early charcoal administration may be useful in partially

preventing the absorption of aripiprazole. Administration of 50 g of activated charcoal, one hour after a single 15-mg oral dose of aripiprazole, decreased the mean AUC and C_{max} of aripiprazole by 50%.

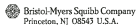
Hemodialysis: Although there is no information on the effect of hemodialysis in treating an overdose with

aripiprazole, hemodialysis is unlikely to be useful in overdose management since aripiprazole is highly bound to plasma proteins.

Tablets manufactured by Otsuka Pharmaceutical Co., Ltd., Tokyo, 101-8535 Japan or Bristol-Myers Squibb Company, Princeton, NJ 08543 USA

Orally disintegrating tablets manufactured by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA Oral Solution manufactured by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA Distributed and marketed by Otsuka America Pharmaceutical, Inc., Rockville, MD 20850 USA Marketed by Bristol-Myers Squibb Company, Princeton, NJ 08543 USA US Patent Nos: 5,006,528 and 6,977,257

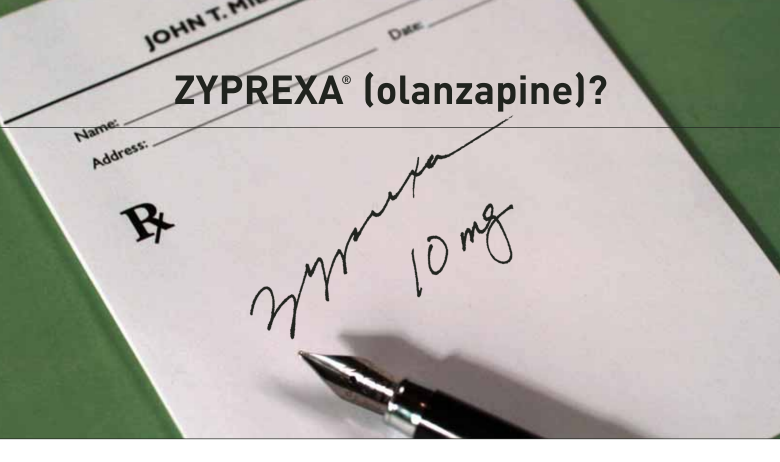
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June 2006



You wrote "ZYPREXA." Will your patient leave the pharmacy with something else?

With over 4,000 drugs on the market and more than 8 million prescriptions filled every day, medication errors can and do occur. For example, ZYPREXA and Zyrtec* (cetirizine HCl) have been mistaken, one for the other, in the past.

To help avoid such medication errors, the Institute for Safe Medication Practices (ISMP) recommends that physicians:

- $\boldsymbol{\cdot}$ Print the medication's brand name and generic name on all prescriptions.
- $\boldsymbol{\cdot}$ Include dosage form, strength, and full instructions.
- Pronounce the name for the patient or caregiver, and have them say it back to you.
- Remind the patient to check for anything unusual (eg, capsules instead of the usual tablets) before they leave the pharmacy.

Please take special care when prescribing any medication. Millions of patients and their families are counting on you.

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SENIOR FACULTY POSITION

The University of North Carolina School of Medicine is seeking a senior psychiatrist at the Associate or Full Professor level (fixed-term) for the position of Clinical Director for a new, modern public psychiatric facility which is under construction in nearby Butner, North Carolina. This 432 bed hospital will provide psychiatric services for the central region of the state, and will include vibrant academic, teaching, and research programs in partnership with both the UNC School of Medicine and Duke University Medical Center. The Clinical Director will be responsible for medical leadership and the planning and organization of clinical, research and teaching functions. Candidates must have an M.D. from an accredited university, be Board Certified/Eligible, and able to obtain medical licensure in North Carolina. A strong track record of administrative leadership in a public academic setting is highly desirable.

Candidates should submit a letter of application, current CV, and the names and contact information of three professional references to:

David Rubinow, M.D.
Meymandi Professor and Chair of Psychiatry
Campus Box #7160
University of North Carolina School of Medicine
Chapel Hill, NC 27599-7160

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their work and life We are pleased to announce tremendous opportunities in rapidly expanding, behavioral health organizations in Lafayette, LA and Longview, TX. We are now seeking general psychiatrists and child/adolescent psychiatrists. • Primary focus will be inpatient and outpatient psychiatric services • Highly competitive compensation package • Relocation available · Board eligibility/board certification required For questions or confidential consideration, please contact Susan Legg: slegg@acadiahealthcare.com P: 770.772.4345 www.acadiahealthcare.com Acadia Healthcare ... Providing hope today for a better tomorrow.

ADULT PSYCHIATRIST (BC/BE)

Southern New Hampshire

Outstanding opportunity for an Adult Psychiatrist interested in outpatient and inpatient care. This opportunity allows a Provider to further develop successful psychiatric services and shared call in a 19-bed unit located in a 330-bed acute-care hospital. Position would be an Associate Medical Directorship with the potential of Medical Director after 1 year. The opportunity offers an extremely competitive salary with outpatient billing incentive.

Opportunity is located in Manchester, New Hampshire, one hour from Boston, the seacoast and the beautiful White Mountains. Affordable housing with no state income or sales tax.

For additional information and to submit CV for consideration, please contact:

Paul Rodden phone: 603-663-8001; fax: 603-663-6441 email: prodden@cmc-nh.org

CHAIR, DEPARTMENT OF PSYCHIATRY Yale University School of Medicine and CHIEF, PSYCHIATRY SERVICE Yale-New Haven Hospital

Yale University announces a search for Chair of the Department of Psychiatry, which presently includes 181 full-time faculty, 70 residents, and more than 60 pre- and postdoctoral trainees at its four research and educational sites. The Department has a distinguished record of success in research, with 12 program project and center grants as well as 8 training grants among its 8 research divisions. The candidate will also serve as Chief of the Psychiatry Service at Yale-New Haven Hospital, which operates the 72-bed Yale-New Haven Psychiatric Hospital, as well as several outnatient sites

The successful candidate will possess an outstanding academic record in clinical, translational, or basic research relevant to the discipline and should be an excellent clinician and educator as well as an experienced administrator with highly developed communication and interpersonal skills. Candidates are expected to meet the requirements for appointment to the senior faculty ranks

Interested applicants should send a curriculum vitae and bibliography, a brief statement of programmatic goals, and a list of professional references prior to February 1, 2007, either electronically (lisa.delizio@yale.edu) or by mail, to:

Margaret K. Hostetter, M.D. c/o Lisa M. DeLizio Department of Pediatrics Yale University School of Medicine 333 Cedar Street, 4086 LMP P.O. Box 208064 New Haven, CT 06520-8064 Telephone: 203-785-6053

Yale University is an equal opportunity, affirmative action employer.

Minorities and women are encouraged to apply.

The Louis A. Johnson VA Medical Center, Clarksburg, West Virginia, is seeking a full-time, permanent, BC/BE psychiatrist. US citizen preferred.

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Send CV to: Physician Recruitment, HPMG, Dole Cannery Square, 501 Alakawa Street, Suite 255, Honolulu, Hawaii 96817-5764, Fax (808) 432-4620 or email: Janice.Omori@kp.org.

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Psychiatrist

Working in Social and Behavioral Health Services, conduct biopsychosocial assessments of pain management program patients and develop comprehensive interdisciplinary treatment plans in coordination with the primary care provider and patient. Requires experience in a full range of psychosocial interventions, techniques, and services to enhance or serve as adjunct therapy to medication for chronic pain. Requires a current, active and unrestricted state license to practice as a medical physician (psychiatrist), US citizenship and proficiency in written and spoken English.

Anchorage offers plenty of opportunity to take in salmon fishing, the arts, or the simplicity of peace and natural serenity. We offer comprehensive benefits, including generous leave, tuition reimbursement, a 24 percent tax deferred cost of living adjustment, and Alaska has no sales or income tax. For a complete job description and application information, please visit www.usajobs.opm.gov, or call Robert Jerdan, HR Specialist, at: (907) 257-5453. To apply, please send your completed application to: Human Resources, AVAHSRO, 2925 DeBarr Road, Anchorage, AK 99508.



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St. John's Clinic is seeking energetic board certified/eligible Child & Adolescent and Adult Psychiatrists to join their well-established, busy Psychiatry Department, in lovely Springfield, Missouri. Inpatient and outpatient practice with large referral base. The inpatient unit is conveniently located close to the physician office building. The department is part of St. John's Clinic, a progressive and growing multispecialty clinic of 470 + physicians in an integrated health care delivery system. For more information about St. John's Health System, please visit www.stjohns.com. St. John's was recently ranked among the TOP 10 in patient satisfaction.

Position offers a competitive salary guarantee. In addition, there is bonus potential and a full array of benefits including health, dental, vision, life and disability insurance, malpractice, CME, vacation, retirement, and relocation.

SPRINGFIELD, MISSOURI, (pop. 200,000) is a growing, sophisticated community. It is home to Missouri's second largest university and a regional home to the arts (symphony, ballet, and theater) and NCAA and semi-professional sports teams (football, basketball, baseball, and hockey). *Employment Review* has named Springfield one of the 10 "Best Places to Live and Work" in the U.S. For more information about Springfield, go to www.springfieldmo.org. EOE/AA Employer.

For more information, contact:

Julie Oliver, Physician Recruiter St. John's Clinic 1965 S. Fremont, Suite 320 Springfield, MO 65804 Phone: (800) 218-5079; Fax: (888) 290-8300 jaoliver@sprg.mercy.net



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DIRECTOR - CENTER FOR ALZHEIMER DISEASE AND RELATED DISORDERS (CADRD)

Southern Illinois University School Of Medicine (SIUSM) is seeking an accomplished clinician scientist to become Director of our Center for Alzheimer Disease and Related Disorders (CADRD), consistent with the mission of SIUSM to become a leading research institution. This tenured/tenure track position will also hold an endowed professorship in Alzheimer disease research.

The CADRD is funded by a large annual grant from the State of Illinois and has a history of continued funding for more than 17 years. The Center has an outreach network of 26 hospitals or clinics throughout Illinois, a brain bank, a neuropsychology program, and specialty clinics for the treatment of Alzheimer disease and other degenerative dementias. Currently, five faculty members are actively engaged in Alzheimer research, with one additional open position. In addition, opportunities are available for collaborations with the Geriatric Center of Excellence, the Department of Psychiatry, and basic science departments such as Pharmacology.

The Director of the SIU CADRD reports to the Dean of SIUSM (institutional and administrative matters) and to the Chair of an appropriate Department (departmental and academic matters). The Director will lead the CADRD, holding responsibility for budget decisions, research direction, staffing, and interactions with other departments, institutes, and agencies. Responsibilities also include working with the Illinois legislature and Departments of Public Health, Aging, Education and Public Aid to develop strategies, external support, and collaborative efforts that will advance the aims of the CADRD in the areas of outreach, teaching, research, and clinical service.

Applicants must be board-certified in a specialty relevant to the activities of SIU CADRD and maintain an active externally-funded research program focused on dementia. Illinois licensure is required prior to employment. Deadline to apply is March 31, 2007 or until filled. Interested applicants should send their CV by mail or email to:

Carol Forestier
Secretary for Dr. Leonard Rybak, MD, Ph.D.
Chairman of Search Committee
SIU School of Medicine
P.O. Box 19643
Springfield, IL 62794-9643
cforestier@siumed.edu.

SIU School of Medicine is an EO/AA Employer.

HEAD, DEPARTMENT OF PSYCHIATRY, FACULTY OF MEDICINE REGIONAL CLINICAL DEPARTMENT HEAD, PSYCHIATRY, CALGARY HEALTH REGION

The University of Calgary, Faculty of Medicine, and the Calgary Health Region invite applications and nominations for the joint position of Head, Department of Psychiatry, Faculty of Medicine, University of Calgary and Regional Clinical Department Head, Psychiatry, Calgary Health Region.

The Department of Psychiatry is part of the rapidly growing Faculty of Medicine, which is in the process of building a major new research facility. The Department has a fully accredited residency training program based throughout all the acute care hospitals and ambulatory facilities of the Calgary Health Region; an active postgraduate and graduate studies program: and a record of accomplishment in research.

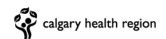
The Regional Clinical Department Head (RCDH), Psychiatry will lead and participate in the development of system-wide clinical services throughout a large and diverse region that incorporates the rapidly growing urban centre of Calgary, as well as surrounding suburban and rural communities. Currently, the Department of Psychiatry has over 120 members in eight clinical divisions. The RCDH will maintain and recruit an appropriately trained physician workforce, ensure high standards of clinical care and ethical conduct of the medical staff, foster maintenance of competence and promote a learning environment for physicians, students, staff and researchers within the Region.

We are searching for an outstanding leader to be the joint RCDH, Psychiatry, and the University's Department Head of Psychiatry. The selected candidate must have proven leadership ability and clinical skills, a strong academic background with demonstrated research and educational achievements, and must be eligible for licensure in the Province of Alberta.

Applications and nominations, including a curriculum vitae, a statement of research interests, leadership philosophy, academic goals, and the names of three referees should be forwarded, by December 31, 2006, to:

D. Grant Gall, MD, FRCPC
Dean, Faculty of Medicine
3330 Hospital Drive, N.W.
Calgary, Alberta, Canada T2N 4N1





In accordance with Canadian Immigration requirements, priority will be given to Canadian citizens and permanent residents of Canada.

The University of Calgary and the Calgary Health Region respect, appreciate and encourage diversity.

LEAD PSYCHIATRIST

The Stratton VA Medical Center located in Albany, New York is recruiting board-certified lead psychiatrist to work in the Behavioral Health Care Line. Position would include supervising professional psychiatric staff as well as clinical responsibilities to be determined.

Interested applicants need to submit current curriculum vitae, license, proof of board certification in Psychiatry to the: Department of Veterans Affairs, Stratton VA Medical Center, HRMS, 113 Holland Avenue, Albany, NY 12208.

For additional information on this position please call Glenn Gilbert: (518) 626-5331.



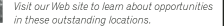
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www.providence.org/physicianopportunities

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CHAIR – Department of Psychiatry and Behavioral Sciences

The Medical University of South Carolina College of Medicine invites applications and nominations for the position of **PROFESSOR and CHAIR** of the Department of Psychiatry. Applicants must be board certified by the American Board of Psychiatry and Neurology, have administrative experience and strong leadership skills, have an outstanding record of accomplishment in research, desire for academic excellence, and have interest in all aspects of teaching and patient care. Particularly important will be a vision of interdisciplinary research programs within and beyond the Medical University of South Carolina.

The Department of Psychiatry is identified as one of the nation's top psychiatric departments with research programs in depression disorders, anxiety, schizophrenia, drug and alcohol addiction and others. For FY 2006, the Department received more than 25 million dollars in research funding and is ranked within the top 20 departments in NIH funding.

The Department is staffed with more than 100 full-time faculty and ample support personnel. Dedicated to educating future clinicians, the Department currently has 80 residents, psychology interns, and fellows enrolled in its graduate education programs.

One of the Department's nationally leading programs is the Center for Drug and Alcohol Programs (CDAP), one of the 14 prestigious centers funded by the National Institute on Alcohol Abuse and Alcoholism (NIAAA) and recognized as a premier program for the treatment of alcohol and other substance abuse disorders. Other Departmental programs include the Brain Stimulation Laboratory, Weight Management Center, an Eating Disorders Program, Psychopharmacology Division, Family Service Research Center, National Crime Victims Center, Clinical Neuroscience Division (CND), and the Clinical Neuropharmacology Program.

The Department's main facility, the Institute of Psychiatry, is a stand alone state-of-the-art building that encompasses in and outpatient clinical services, research, and educational programs. Fully accredited by JCAHO, the Institute has 90 licensed inpatient beds. Additional clinical sites include the Charleston Veterans Affairs Medical Center (16 bed), and a number of community-based clinics and day treatment centers. Interested persons can send curriculum vitae to: L. Lyndon Key, M.D., Chair, Psychiatry Search Committee, c/o Beverly Carson Dean's Office, College of Medicine, Medical University of South Carolina, 96 Jonathan Lucas Street, PO Box 250617, Charleston, SC 29425 or Email: carsonb@musc.edu Deadline for application is January 15, 2007.

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If you would like to learn more about the area, please view these web sites:

www.fcvb.org/html/kalispell.html www.fcvb.org

www.kalispellchamber.com www.nps.gov/glac/photos.htm

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The LSU Health Sciences Center is pleased to announce the availability of the Paul J. Ramsey Chair of Psychiatry. This Chair honors a citizen of Louisiana who dedicated his life to improving the community especially for our city and our state with programs to deal with violence in schools and to help our at risk youths.

We are seeking applicants who have a demonstrated ability to successfully collaborate with diverse community groups and to attract funding not only from various granting agencies but also from private foundations and from other independent sources such as donors within the community. The chair will continue and enrich the community interventions of the Department of Psychiatry and its Paul J. Ramsay Chair through designing, developing and implementing both systems and model demonstration projects in areas such as, programs to reduce the incidence of violence, methods for early intervention in dealing with the impact of exposure to violence and mitigation of the effects it has on the children of New Orleans, as well as the continued commitment to mental health care in the community.

Applicant should possess leadership and administrative skills, and will possess proven scholarship, grantsmanship, community involvement, and respect in the area of mental health of young children necessary to attract future funding for new programs, further develop the teaching and training components of existing programmatic efforts of the department, and continue to address the community's needs in an active and expanding manner.

Applicants must submit comprehensive curriculum vitae and a statement of community interventions interests and the names and contact information of at least three individuals who will serve as references to:

Dr. Howard J. Osofsky, Chair and Professor **LSUHSC** Department of Psychiatry Lions Eye Clinic 2020 Gravier Street, New Orleans, Louisiana 70112

LSUHSC is an EEO/AAE.

POST-DOCTORAL RESEARCH FELLOWSHIP IN CHILD PSYCHIATRY UNIVERSITY OF PITTSBURGH SCHOOL OF MEDICINE

A post-doctoral research fellowship position is available in our clinical research program focusing on the following areas: the longitudinal study of the characteristics, course, risk factors, psychobiology (including functional magnetic resonance imaging) and treatment of childhood/adolescent affective and anxiety disorders, autism, attention deficit, disruptive disorders and eating disorders.

Our Advanced Center for Interventions and Services Research for Early-Onset Mood and Anxiety Disorders and three program projects (in the Psychobiology of Affective & Anxiety Disorders, Behavior Genetics of Affective/Anxiety Disorders, and the Neurobiology of Autism) provide the opportunity to work and study with some of the country's leading clinical researchers. Potential mentors include:

David Axelson, M.D. - Bipolar Disorder Boris Birmaher, M.D. - Mood & Anxiety Disorders David Brent, M.D. - Mood & Anxiety Disorders and Suicidal Behavior Oscar Bukstein, M.D., M.P.H. - Adolescent Substance Bernie Devlin, Ph.D. - Psuchiatric Genetics Walter Kaye, M.D. - Eating Disorders

David Kolko, Ph.D. - Conduct Disorders Rolf Loeber, Ph.D. – Conduct Disorders Beatriz Luna, Ph.D. - fMRI Nancy Minshew, M.D. - Autism Brooke Molina, Ph.D. - ADHD James Perel, Ph.D. - Psychopharmacology Neal Ryan, M.D. - Mood & Anxiety Disorder

We seek individuals either with an M.D. who have completed an accredited residency program in general and/or child psychiatry or a Ph.D. in psychology (clinical /quantitative) from an APA-accredited program with a broad-based intellectual background, evident potential for academic/psychiatric research, and ability to think creatively. Must have clinical experience in psychiatric inpatient/ outpatient and/or pediatric setting, familiarity with psychiatric nosology and interest in basic or applied research in developmental psychopathology. A high proportion of our graduates have received external funding.

Please submit vita and three letters of recommendation to:

David A. Brent, M.D. Professor of Psychiatry, Pediatrics and Epidemiology University of Pittsburgh School of Medicine Western Psychiatric Institute and Clinic 3811 O'Hara Street, BT 313 Pittsburgh, PA 15213 Email: brentda@upmc.edu FAX: (412) 246-5344

THE NOVARTIS CHAIR IN SCHIZOPHRENIA RESEARCH



The **Faculty of Medicine** invites applications for a full-time academic position as The Novartis Chair in Schizophrenia Research. The successful candidate will be a clinical or basic scientist with a distinguished reputation and proven leadership in research, education and knowledge translation in the field of schizophrenia. The incumbent will provide leadership in the development and maintenance of a world-class schizophrenia research program, will demonstrate a commitment to excellence in teaching and patient care, if applicable, as well as attract outstanding students, trainees and scholars in this field. If the research is patient-based and involves the provision of patient care, the selected individual must be eligible for licensure in the Province of Alberta. The Chair holder may be appointed in the Department of Psychiatry and other departments, as appropriate, and may become a full member of the Hotchkiss Brain Institute.

The successful candidate's research interests may fall within any of the four CIHR pillars; basic biomedical, clinical, health services, and/or population health research. The selected candidate may compete for salary support and establishment funding from the Alberta Heritage Foundation for Medical Research. We anticipate generous start-up funds will be available to a qualified candidate.

With over 400 full-time members, the Faculty of Medicine is a leader in health research with an international reputation for excellence and innovation. A new research facility slated to open in 2007 will allow researchers to investigate scientific questions collaboratively in a unique setting that facilitates multidisciplinary studies with state-of-the-art investigative tools. Calgary is a vibrant, multicultural city of 1,000,000 near the Rocky Mountains, Banff National Park and Lake Louise.

The Department of Psychiatry and the Hotchkiss Brain Institute are working together to enable integrated, outcome-focused research and education activities. One of the eight Hotchkiss Brain Institute research programs is focused on population mental health research and education activities. See the website www.hbi.ucalgary.ca for more information on the institute.

Please submit curriculum vitae, a statement of research interests and goals, and the names of three referees by **December 31, 2006**, to:

Dr. P.A. Sokol

Vice-Dean, Faculty of Medicine 3330 Hospital Drive N.W. Calgary, Alberta, Canada T2N 4N1

In accordance with Canadian immigration requirements, priority will be given to Canadian citizens and permanent residents of Canada. The University of Calgary respects, appreciates and encourages diversity.

www.ucalgary.ca

MINORITY RESEARCH TRAINING IN PSYCHIATRY

The Program for Minority Research Training in Psychiatry (PMRTP) is funded by the National Institute of Mental Health (NIMH). Through it, the American Psychiatric Institute of Research and Education (APIRE) sponsors training of students. medical minority psychiatric residents, and fellows who are interested in research by providing advice, placement assistance, tuition, stipends, travel and other expenses. The annual application deadline for research fellows is December 1 and for summer medical students is April 1. The director of the program is Darrel A. Regier, M.D., M.P.H.; the project manager is Ernesto A. Guerra.

For more information:

Call: 1-800-852-1390 or 703-907-8622
E-mail: eguerra@psych.org
Write: PMRTP
American Psychiatric Institute for
Research and Education
1000 Wilson Blvd, Ste. 1825
Arlington, VA 22209-3901

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Reference: 1. Data on file, Wyeth Pharmaceuticals Inc.

Please see brief summary of Prescribing Information on adjacent pages.

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BRIEF SUMMARY. See package insert for full prescribing information.

Suicidality in Children and Adolescents

Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with Major Depressive Disorder (MDD) and other psychiatric disorders. Anyone considering the use of EFFEXOR XR or any other antidepressant in a child or adolescent must balance this risk with the clinical need. Patients who are started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. EFFEXOR XR is not approved for use in pediatric patients. (See Warnings and Precautions: Pediatric Use.)

Pooled analyses of short-term (4 to 16 weeks) placebo-controlled trials of 9 antidepressant drugs (SSRIs and others) in children and adolescents with Major Depressive Disorder (MDD), obsessive-compulsive disorder (OCD), or other psychiatric disorders (a total of 24 trials involving over 4,400 patients) have revealed a greater risk of adverse events representing suicidal thinking or behavior (suicidality) during the first few months of treatment in those receiving antidepressants. The average risk of such events in patients receiving antidepressants was 4%, twice the placebo risk of 2%. No suicides occurred in these trials suicides occurred in these trials.

CONTRAINDICATIONS: Hypersensitivity to venlafaxine hydrochloride or to any excipients in the formulation. Concomitant use in patients taking monoamine oxidase inhibitors (MAOIs). WARNINGS: Clinical Worsening and Suicide Risk—Patients with major depressive disorder (MDD), both adult and pediatric, may experience Concomitant use in patients taking monoamine oxidase inhibitors (MAOIs). WARNINGS: Clinical Worsening and Suicide Risk.—Patients with major depressive disorder (MDD), both adult and pediatric, may experience worsening of their depression and/or the emergence of suicidal ideation and behavior (suicidality) or unusual changes in behavior, whether or not they are taking antidepressant medications, and this risk may persist until significant remission occurs. There has been a long-standing concern that antidepressants may have a role in inducing worsening of depression and the emergence of suicidality in certain patients. Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with MDD and other psychiatric disorders. It is unknown whether the suicidality risk extends to adults. All pediatric patients being treated with antidepressants for any indication should be observed closely for clinical worsening, suicidality, and unusual changes in behavior, especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases. Anxiety, agitation, panic attacks, insomnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomania, and mania have been reported in adult and pediatric patients being treated with antidepressants for MDD and other indications, both psychiatric and nonpsychiatric. Although a causal link between the emergence of such symptoms and either the worsening of depression and/or the emergence of suicidal impulses has not been established, there is concern that such symptoms and either the vacicidality, or symptoms may represent precursors to emerging suicidality. Consideration, in patients whose depression is persistently worse, or who are experiencing emergent suicidality or symptoms that might be precursors to oversening depression or solicidality as is feasibly discontinuing the medication, in patients whose depression is pers agitation, irritability, unusual changes in behavior, and the other symptoms described above, as well as the emergence of suicidality, and to report such symptoms immediately to health care providers. Such monitoring should include daily observation by families and caregivers. Prescriptions for Effexor XR should be written for the smallest quantity of capsules consistent with good patient management, in order to reduce the risk of overdose. Families and caregivers of adults being treated for depression should be similarly advised. Screening Patients for Bipolar Disorder. A major depressive episode may be the initial presentation of bipolar disorder. It is generally believed that treating such an episode with an antidepressant alone may increase the likelihood of precipitation of a mixed/manic episode in patients at risk for bipolar disorder. Whether any of the symptoms described above represent such a conversion is unknown. Prior to initiating antidepressant treatment, patients with depressive symptoms should be screened to determine if they are at risk for bipolar disorder; such screening should include a detailed psychiatric history, including a family history of suicide, bipolar disorder, and depression. Effexor XR is not approved for use in treating bipolar depression. Potential for Interaction with MAOIs—Adverse reactions, some serious, have been reported in patients who recently discontinued an MAOI and started on venlafaxine, or who recently discontinued venlafaxine prior to initiation of an MAOI. These reactions included tremor, myoclonus, diaphoresis, nausea, vomiting, flushing, dizziness, hyperthermia with features resembling neuroleptic malignant syndrome, seizures, and death. Effexor XR should not be used in combination with an MAOI, or within at least 14 days of discontinuing treatment with an MAOI. At least 7 days should be allowed after stopping venlafaxine before startling an MAOI. Secrotonin Syndrome—The development of potentially life-threatening serotonin syndrome may occur with Effexor XR t had ≥7% loss of body weight and no patients discontinued for weight loss. In 12-week PD trials, 3% of Effexor XR patients had ≥7% loss of body weight, and no patients discontinued for weight loss. The safety and efficacy of ventafaxine in combination with weight loss agents, including phentermine, have not been established. Coadministration of Effexor XR and weight loss agents is not recommended. Effexor XR is not indicated for weight loss alone or in combination with other products. Pediatric Patients. Weight loss was seen in patients aged 6-17 receiving Effexor XR. More Effexor XR patients than placebo patients experienced weight loss was seen in patients aged 6-17 receiving Effexor XR. More Effexor XR patients w. 3.6% of placebo patients; Pc-0.001) and the SAD study (47% of Effexor XR patients vs. 14% of placebo patients; Pc-0.001). Weight loss was not limited to patients with treatment-emergent ancreax (decreased appetite). Children and adolescents in a 6-month MDD study had increases in weight less than expected based on data from age—and sex-matched peers. The difference between observed and expected weight rain was length referred free in Mediate. in weight less than expected based on data from age- and sex-matched peers. The difference between observed and expected weight gain was larger for children -12 years old than for adolescents ≥12 years old ... Changes in Helght. Pediatric Patients: In 8-week GAD studies, Effexor XR patients aged 6-17 grew an average of 0.3 cm (n=122), while placebo patients grew an average of 1.0 cm (n=132); P=0.041. This difference in height increase was most notable in patients <12. In 8-week MDD studies, Effexor XR patients grew an average of 0.8 cm (n=146), while placebo patients grew an average of 0.7 cm (n=147). During the 16-week placebo-controlled SAD study, both the Effexor XR (n=109) and the placebo (n=112) patients grew an average of 1.0 cm. In the 6-month MDD study, children and adolescents had height increases less than expected based on data from age- and sex-matched peers. The difference between observed and expected growth rates was larger for children -12 years old than for adolescents =12 years old. Changes in Appetite: Adult Patients: Treatment-emergent anorexia was more commonly reported for

Effexor XR (8%) than placebo (4%) patients in MDD studies. The discontinuation rate for anorexia was 1.0% in MDD studies. Treatment-emergent anorexia was more commonly reported for Effexor XR (8%) than placebo (2%) patients in GAD studies. The discontinuation rate for anorexia was 0.9% for up to 8 weeks in GAD studies. Treatment-emergent anorexia was more commonly reported for Effexor XR (20%) than placebo (2%) patients in SAD studies. The discontinuation rate for anorexia was 0.4% for up to 12 weeks in SAD studies. Treatment-emergent anorexia was more commonly reported for Effexor XR (8%) than placebo (3%) patients in PD studies. The discontinuation rate for anorexia was 0.4% for Effexor XR, patients in 12-week PD studies. **Pediatric Patients:** Decreased appetite was seen in pediatric patients receiving Effexor XR, in GAD and MDD trials, 10% of Effexor XR patients aged 6-17 for up to 8 weeks and 3% of placebo patients had treatment-emergent anorexia. None of the patients receiving Effexor XR and placebo-controlled trial for SAD, 22% and 3% of patients aged 8-17 treated for up to 16 weeks with Effexor XR and placebo, respectively, reported treatment-emergent anorexia (decreased appetite). The discontinuation rates for anorexia were 0.7% and 0.0% for patients receiving Effexor XR and placebo, respectively, by fatients receiving effector XR and placebo. (decreased appetite). The discontinuation rates for anorexia were 0.7% and 0.0% for patients receiving Effexor XR and placebo, respectively; the discontinuation rates for weight loss were 0.7% for patients receiving either Effexor XR or placebo. Activation of Mania/Hypomania* Mania or hypomania has occurred during short-term depression and PD studies. As with all drugs effective in the treatment of MDD, Effexor XR should be used cautiously in patients with a history of mania. Hyponatremia: Hyponatremia and/or the syndrome of inappropriate antidiuretic hormone secretion (SIADH) may occur with venlafaxine. Consider this in patients who are volume-depleted, elderly, or taking diuretics. Seizures: Nal II premarketing depression trials with Effexor, seizures were reported in 0.3% of venlafaxine patients. Use cautiously in patients with a history of seizures. Discontinue in any patient who develops seizures. Abnormal Bleeding: Ahonormal bleeding (most commonly ecctymosis) has been reported. Serum Cholesterol Elevation. Clinically relevant increases in serum cholesterol were seen in 5.3% of venlafaxine patients and 0.0% of placebo patients treated for at least 3 months in trials. Consider measurement of serum cholesterol levels during long-term treatment. *Use in Patients With Concomitant Illness*: Use Effexor XR cautiously in patients with diseases or conditions that could affect hemodynamic responses or metabolism. Ventadaxine has not been evaluated in patients with recent history of MI or unstable heart disease. Increases in QT interval (QTc) have been reported in clinical studies. Exercise caution in patients whose underlying medical conditions might be compromised by increases in heart rate. In patients with renal impairment or cirrhosis of the liver, the clearances of venlataxine and its active metabolites were decreased, prolonging the elimination half-likes. A lower dose may be necessary, use with caution in such patients. Information for Patients—Prescribers or other health professionals should inform patients, their families, and their caregivers about the benefits and risks associated with treatment with Effexor XR and should counsel them in its appropriate use. A patient Medication Guide About Using Antidepressants in Children and Teenagers is available for Efferor XR. The prescriber or health professional should instruct patients, their families, and their caregivers to read the Medication Guide and should assist them in understanding its contents. Patients should be given the opportunity the Medication Guide and should assist them in understanding its contents. Patients should be given the opportunity to discuss the contents of the Medication Guide and to obtain answers to any questions they may have. The complete text of the Medication Guide is available at www.new.engex.or.com or in the approved prescribing information. Patients should be advised of the following issues and asked to alert their prescriber if these occur while taking Effexor XR. Clinical Worsening and Suicide Risk: Patients, their families, and their caregivers should be encouraged to be alert to the emergence of symptoms listed in WARNINGS. Clinical Worsening and Suicide Risk, especially those seen early during antidepressant treatment and when the dose is adjusted up or down. Families and caregivers of patients should be advised to observe for the emergence of such symptoms on a day-to-day basis, since changes may be abrupt. Such symptoms should be reported to the patient's prescriber or health professional, especially if they are severe, abrupt in onset, or were not part of the patient's presenting symptoms. Symptoms such as these may be associated with an increased risk for suicidal thinking and behavior and indicate a need for very close monitoring and possibly changes in the medication. Caution patients 1) about operating hazardous machinery, including automobiles, until they are reasonably sure that ventafaxine does not adversely affect their abilities; 2) to avoid alcohol while taking Effexor XR; and 3) about the risk of serotonin syndrome with the concomitant use of Effexor XR and triptans, tramadof, thyptophana supplements, or other serotonergic agents. Patients should be advised to notify their physician 1) if they Effexor XR, and 3) about the risk of serotonin syndrome with the concomitant use of Effexor XR and triptans, tramadol, tryptophan supplements, or other serotonergic agents. Patients should be advised to notify their physician 1) if they become pregnant or intend to become pregnant during therapy, or if they are nursing; 2) about other prescription or over-the-counter drugs, including herbal preparations and nutritional supplements they are taking or plan to take; 3) if they develop a rash, hives, or related allergic phenomens; or 4) if they have a history of glaucoma or increased intraocular pressure. Laboratory Tests—No specific laboratory tests are recommended. Drug Interactions—Alcohof. A single dose of ethanol had no effect on the pharmacokinetics (PK) of ventilataine or O-desmethylyvenializatine (DV), and ventilatatione that of the psychomotor and psychometric effects induced by ethanol. Climetidine. Use caution when administering ventalaxine with cimetidine to patients with pre-existing hypertension or hepatic visfunction, and the elderly Diazepam. A single dose of diazepam did not appear to affect the PK of either ventalaxine or ObV Ventalaxine during the propertion of the psychomotor and psychometric effects induced by diazepam. A laboratory tests are recommended to the psychomotor and psychometric effects induced by diazepam. A laboratory the psychomotor and psychometric effects induced by diazepam did not appear to affect the PK of either ventalaxine or ODV ventalaxine decreased total oral-dose clearance of halopendol, resulting in a 70% increase in halopendol AUC. The haloperidol C_{max} increased 8%, but the haloperidol elimination half-life was unchanged. Lithium A single dose of lithium did not appear to affect the PK of either ventalaxine or ODV. Ventalaxine had no effect on the PK of lithium. Drugs Highly Bound to Plasma Proteins: Ventalaxine is not highly bound to plasma proteins; coadministration of Effexor XR with a highly protein-bound drug should not cause increased free concentrations of th CyP2D6, Drugs inhibiting this isoenzymes. CYP2D6 Inhibitors: Venlataxine is metabolized to its active metabolite, ODV, by CYP2D6. Drugs inhibiting this isoenzyme have the potential to increase plasma concentrations of venlataxine and decrease concentrations of ODV. No dosage adjustment is required when venlataxine is coadministered with a CYP2D6 inhibitor. Concomitant use of venlataxine with drug treatment(s) that potentially inhibits both CYP2D6 inhibitor. Concomitant uses of venlataxine with drug treatment(s) that potentially inhibits both CYP2D6 inhibitor. Concomitant uses of venlataxine, has not been studied. Use caution if therapy includes venlataxine and any agent(s) that produces simultaneous inhibition of these two enzyme systems. Drugs Metabolized by Cytochrome P450 Isoenzymes. Venlataxine is a relatively weak inhibitor of CYP2D6. Venlataxine did not inhibit CYP1A2 and CYP3A4, CYP2D9 (in vitro), or CYP2C19. Impramine Venlataxine did not affect the PK of enplataxine. In the Pking of interparatine and 2-OH-imipramine AUCs increased by 2-5-4.5 fold. Imipramine did not affect the PK of venlataxine sightly inhibited the CYP2D6-mediated metabolism of risperidone to its active metabolite, 9-hydroxyrisperidone, resulting in a ~32% increase in risperidone AUC. Venlataxine coadministration did not significantly alter the PK profile of the total active moiety (risperidone plus 9-hydroxyrisperidone). CYP3A4: Venlataxine add not ninhibit CYP3A4 in vitro and in vivo. of Shafimavin in a study of 9 healthy volunteers, venlataxine administration did not affect the PK of venlataxine and ODV. CYP1A2: Venlataxine did not ninhibit CYP2A9 in vitro and in vivo. CYP2C9-mediated formation of 4-hydroxy-foblutamide. CYP2C9: Venlataxine did not ninhibit the metabolism of diazepam, which is partiality metabolized by CYP2C19 (see Diazepam about a scaled on the metabolism of diazepam, which is partiality metabolized by CYP2C19 (see Diazepam about a scaled on the metabolism of diazepam, which is partiality metabolized by CYP2C19 (see Venlafaxine and DDV were not mutagenic in the Ames reverse mutation assay in Salmonella bacteria or the CHO/HGPRT mammalian cell forward gene mutation assay. Venlafaxine was not clastogenic in several assays. DDV elicited a clastogenic response in the in vivo chromosomal aberration assay in rat bone marrow. Impairment of Fertility. No effects on reproduction or fertility in rats were noted at oral doses of up to 2 times the MRHD on a mg/m² basis, **Pregnancy**—*Teratogenic Effects*—*Pregnancy Category C.* Reproduction studies in rats given 2.5 times, and rabbits given 4 times the MRHD (mg/m² basis) revealed no malformations in offspring. However, in rats given 2.5 times the MRHD, there was a decrease in pup weight, an increase in stillborn pups, and an increase in pup deaths during the first 5 days of lactation when dosing began during pregnancy and continued until weaning. There are no adequate and well-controlled studies in pregnant women; use Effexor XR during pregnancy only if clearly needed. Monteratogenic Effects. Neonates exposed to Effexor XR late in the third trimester have developed complications requiring prolonged hospitalization, respiratory support, and tube feeding. Complications can arise immediately upon requiring prolonged hospitalization, respiratory support, and tube feeding. Complications can arise immediately upon delivery. Reports include respiratory distress, cyanosis, apnea, sezures, temperature instability, feeding difficulty, womiting, hypoglycemia, hypotonia, hypertonia, adequately assessed the impact of Effexor XR on growth, development, and maturation of children and adolescents. Studies suggest Effexor XR may adversely affect weight and height (see PRECAUTIONS-General, Changes in Height and Changes in Weight). Should the decision be made to treat a pediatric patient with Effexor XR, regular monitoring of weight and height is recommended during treatment, particularly if long term. The safety of Effexor XR for pediatric patients has not been assessed for chronic treatment >6 months. In studies in patients aged 6-17, blood pressure

and cholesterol increases considered to be clinically relevant were similar to that observed in adult patients. The precautions for adults apply to pediatric patients. Geriatric Use—No overall differences in effectiveness or safety were observed between geriatric and younger patients. Greater sensitivity of some older individuals cannot be ruled out. Hyponatremia and SIADH have been reported, usually in the elderly. AVUENSE REACTIONS: Associated with Discontinuation of Treatment—The most common events leading to discontinuation in MDD, GAD, SAD, and PD trials included nausea, anorexia, anaively, impotence, dry mouth, dizziness, insomnia, somnolence, hypertension, darrhea, paresthesia, termor, abnormal (mostly blurred) vision, abnormal (mostly delayed) ejaculation, asthenia, vomiting, nervousness, headache, vasodilatation, thinking abnormal, decreased libido, and sweating. Commonly Observed Adverse Events in Controlled Clinical Trials for MDD, GAD, SAD, and PD—Body as a Whole; asthenia, headache, flu syndrome, accidental injury, abdominal pain. Cardiovascular: vasodilatation, hypertension, palpitation. Digestive: nausea, constipation, anorexia, vomiting, fatulence, diarrhea, enuctation, Metabolic/Nutritional: weight loss. Mervous System: disziness, somnolence, insomnia, dry mouth, nervousness, abnormal direams, tremor, depression, hypertonia, paresthesia, libido decreased, agitation, anxiety, twitching. Respiratory System: pharyngitis, yawn, sinusitis. Skin: sweating, Special Senses: anormal vision. Uroqenital System: abnormal ejaculation, impotence, orgasmic dysfunction (including anorgasmia) in females. Vital Sign Changes: Effexor XR was associated with a mean increase in pulse rate of about 2 beats/min in depression and GAD trials and a mean increase in pulse rate of increase in pulse rate of about 2 beats/min in depression and GAD trials and a mean increase in pulse rate of about 2 beats/min in depression and GAD trials and a mean increase in pulse rate of 4 beats/min in SAD trials. (See WARNINGS-Sustained Hypertension). Laboratory Changes: Clinically relevant increases in serum cholesterol were noted in Effexor XR clinical trials. Increases were duration dependent over the study period and tended to be greater with higher doses. Other Events Observed During the Premarketing Evaluation of Effexor and Effexor XR—N=6,670. "Frequent"-events occurring in at least 1/100 patients, "infrequent"=1/100 to 1/1000 patients, "infrequent"=1/100 to 1/1000 patients, "infrequent"=1/100 to 1/1000 patients, "infrequent"=1/100 patients, "infrequent"=1/100 patients, "infrequent"=1/100 patients, "infrequent" intentional injury, malaise, monificialis, neck rigidity, pelvic pain, photosensitivity reaction, suicide attempt, withdrawal syndrome; Rare: appendicitis, bacteremia, carcinoma, carluitis. Cardiovascular system - Frequent increase dispetite, entending and perioditis, gastroitis, gastroitis, gastroitis, gastroitis, gastroitis, gastroitis, pastroitis, periodioritis, proditis, rectal disorder, sopphagel perioditis, perioditis, proditis, rectal disorder, salvary gland enlargement, increased appetite; infrequent brusism, oblist, prio, richillis, cholecystitis, choletitissis, esophageal spasms, duodentiis, hematemesis, gastrosophageal reflux disease, gastroitis, periodioritis, periodioritis, proditis, rectal disorder, salvary gland enlargement, increased aboration, soft stools, tongue cholecystitis, choleithiasis, esophageal spasms, duodentits, hematemesis, gastroesophageal reflux disease, gastrointestinal hemorrhage, gum hemorrhage, hepatitis, ielitis, jaundice, intestinal obstruction, liver tenderness, parotitis, periodintitis, proctitis, rectal disorder, salivary gland enlargement, increased salivation, soft stools, tongue discoloration. Endocrine. system - Fare: galactorrhoea, goiter, hyperthyroidism, hypotryloidism, thyroid nodule, hypotralicy, thrombocythemia; Rare: basophilia, bleeding time increased, cyanosis, eosinophilia, lymphocytosis, multiple myeloma, purpura, thrombocytopenia. Metabolic and nutritional- Frequent edema, weight gain; Infrequent: alkaline phosphatase increased, dehydration, hypercholesteremia, hyperdycemia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hypotylogenia, hyperchalemia, hyperphosphatemia, bylogentalemia, bylogentorienemia, ermai. Musculoskeletal system: Frequent: atrihagia; Infrequent: arithmis, arthrosis, bone spurs, bursitis, leg cramps, myasthenia, tenosynovitis; Rare: bone pain, pathological fracture, musculoskeletal system: Frequent arthralgia; Infrequent arthritis, arthrosis, bone spurs, bursitis, leg cramps, myasthenia, tenosynovitis; Rare: bone pain, pathological fracture, musculoskeletal system: Frequent: arthralgia; Infrequent arthritis, arthrosis, bone spurs, bursitis, leg cramps, myasthenia, tenosynovitis; Rare: bone pain, pathological fracture, musculoskeletal system: Frequent: arthralgia; Infrequent arthritis, arthrosis, bone spurs, bursitis, leg cramps, myasthenia, tenosynovitis; Rare: bone pain, pathological fracture, musculoskeletal system: Frequent arthralgia; Infrequent archive, providia de nalis, erythema nodosum, extoliativa dermatitis, lichenoid dermatitis, hair discoloration, skin discoloration, sturniculosis, instruttism, leukoderma, miliaria, petechial rash, pruritic rash, pustular rash, vesiculobilulos rash, seborrhea, skin atrophy, skin hypertrophy, skin striae, sweating decreased. Special senses. - Frequent abnormality of accommodation, prydrasis, taste perversion; infrequent conjunctivitis, diplopid, oft yeyes, eye pain, hyperacusis, otitis media, parosmia, photophobia, taste loss, visual field defect; Rare: blepharitis, cataract, chromatopsia, conjunctival edema, comeal iesion, deafness, exophithalmos, eye hemorrhage, glaucoma, retinal hemorrhage, subconjunctival hemorrhage subconjunctival hemorrhage subconjunctival hemorrhage system - Frequent: prostatic disorder (prostatitis, enlarged prostate, and prostate irritability), urination impaired; infrequent: albuminuria, amenorrhae, oxistitis, dysuria, hematuria, kidney calculus, kidney pain, leukorrhea, menorrhagia, metrorrhagia, nocturia, breast pain, polyuria, puria, urinary incontinence, urinary retention, urinary urgenov, vaginal hemorrhage, vaginitis; Rare: abortion, anuria, balantis; baldeder pain, breast discharge, breast engorgement, breast enlargement, endometriosis, female lactation, fibrocystic breast, calcium crystalluria, cervicitis, orchitis, ovarian cyst, prolonged erection, gynecomastia (male), hypomenorrhea, kidney function abnormal, mastitis, ovarian cyst, prolonged erection, gynecomastia (male), hypomenorrhea, kidney function abnormal, mastitis, norchitis, ovarian cyst, prolonged erection, gynecomastia (male), hypomenorrhea, kidney function abnormal dryness. Postmarketing Reports: agranulocytosis, anaphylaxis, aplastic anemia, catatonia, congenital anomalies, CPK increased, deep vein thromobophiebitis, delimim, EKG abnormalities such as OT prolongation; cardiac arrhythmias including atrial fibrillation, supraventricular tachycardia, venticular extrasystoles, and rare reports of ventricular abnormalities of unspecified l recovered), neutropenia, night sweats, pancreatitis, pancytopenia, panic, prolactin increased, renal failure, rhabdomyolysis, serotonin syndrome, shock-like electrical sensations or timitus (in some cases, subsequent to the discontinuation of venifaxine or tapering of dose), and SIADH (usually in the elderly). Elevated clozapine levels that were temporally associated with adverse events, including seizures, have been reported following the addition of use of initial control of the contro

Depression can recur many times.





Extending the body of evidence

2-YEAR RECURRENCE PREVENTION

data for EFFEXOR XR1

IMPORTANT TREATMENT CONSIDERATIONS

Suicidality in Children and Adolescents

Antidepressants increased the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents with Major Depressive Disorder (MDD) and other psychiatric disorders. Anyone considering the use of EFFEXOR XR or any other antidepressant in a child or adolescent must balance this risk with the clinical need. Patients who are started on therapy should be observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. EFFEXOR XR is not approved for use in pediatric patients.

- EFFEXOR XR is contraindicated in patients taking monoamine oxidase inhibitors (MAOIs).
- Adult and pediatric patients taking antidepressants can experience
 worsening of their depression and/or the emergence of suicidality.
 Patients should be observed closely for clinical worsening
 and suicidality, especially at the beginning of drug therapy,
 or at the time of increases or decreases in dose. Anxiety,
 agitation, panic attacks, insomnia, irritability, hostility, aggressiveness,
 impulsivity, akathisia, hypomania, and mania have been reported and
 may represent precursors to emerging suicidality. Stopping or
 modifying therapy should be considered especially when symptoms
 are severe, abrupt in onset, or not part of presenting symptoms.

- The development of potentially life-threatening serotonin syndrome may occur when EFFEXOR XR is coadministered with other drugs that may affect the serotonergic neurotransmitter systems. Concomitant use of EFFEXOR XR with MAOIs is contraindicated. If concomitant use of EFFEXOR XR with an SSRI, SNRI, or a triptan is clinically warranted, careful observation of the patient is advised. Concomitant use of EFFEXOR XR with tryptophan supplements is not recommended.
- Treatment with venlafaxine is associated with sustained increases in blood pressure (BP) in some patients. Postmarketing cases of elevated BP requiring immediate treatment have been reported. Pre-existing hypertension should be controlled. Regular BP monitoring is recommended.
- Mydriasis has been reported in association with venlafaxine; therefore, patients with raised intraocular pressure or those at risk of acute narrow-angle glaucoma (angle-closure glaucoma) should be monitored.
- Abrupt discontinuation or dose reduction has been associated with discontinuation symptoms. Patients should be counseled on possible discontinuation symptoms and monitored while discontinuing the drug; the dose should be tapered gradually.



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