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February 2007

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Increased Mortality in Elderly Patients with Dementia-Related Psychosis: Elderly patients with dementia-related psychosis treated with adpical 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 hybica 110 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 tax 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.

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 schizoohrenic patients.

schzopnenic patients — *QT Prolongation*: Because 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 mycoardial infarction, or with uncompensated heart failure (see WARNINGS). Pharmacokinetic/pharmacodynamic studies between GEODON and other drugs that prolong the QT interval have beformed. An additive effect of GEODON and other drugs that prolong the QT interval cannot be excluded. Therefore, GEODON should not be given with dofebilde, sotalol, quinkline, other Class Ia and III anti-arrhythmics, mesoridazine, thioridazine, chlorpromazine, droperidol, and all arthythmics with Of prolingation by some either drugs. EECODN's contrained and in patients with a known history of CII ordinary and including companies and propriative, where could be prever with detailed, extending countries, and an advantage of the countries of the co protectin levels in humans. Tissue cultime experiments indicate that approximately one third of human breast cancers are prolacin dependent in vitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancers are prolacin dependent in vitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancers are prolacin dependent in vitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancers are prolacin of the previously detected by the previously detected by

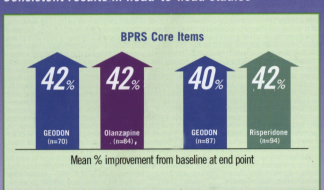
information and instructions in the Patient Information Sectionshould be discussed with patients. Laboratory Tests: Patients being considered for GE000N treatment who are at risk of significant electrolyte disturbances should have baseline serum potassium and magnesium measurements. Low serum potassium and magnesium should be repleted before treatment. Patients who are started on diuretics during GE000N therapy need periodic monitoring of serum potassium and magnesium. Discontinue GE000N in patients who are found to have persistent OT₀ measurements - 500 msec (see WARNINGS). Drug Interactions: (1) GE000N should not be used with any drug that prolongs the OT interval. (2) Given the primary ON. Seffects of GE000N topic of the primary of the primary of the original properties of the primary of the primary of the original properties of the primary of the original properties or the original properties of the original properties or the original properties Showed und execution the aller interdonism of exercimental and reference to the control of the c 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 adenorma and carcinoma, and mammany gland adenorma and actionations, and mammany gland adenorma descentions are descentionated to describe the control of the control of the production and the control of the control und put for commanded that women reaching GEODON should not breast teet. Partialized Use: The settler of GEODON in other careful of human fulls. It is recommended that women reaching GEODON should not breast teet. Partialized Use: The settler and effectiveness of GEODON in predictive patients for the most beautified. General Description of the proportional y 600 per description for the proportional y 600 per description of GEODON in clinical studies. 2 4/9 (1995) were 55 years of age or over. In general there was no indication of any different tolerability for GEODON in clinical studies. 2 4/9 (1995) were 55 years of age or over. In general there was no indication of any different tolerability for GEODON in the middle description of GEODON in the dendry compared to your good of the compared of multiple factors that might in crosses the proportion of GEODON in the dendry compared to your good of the compared of multiple factors that might in crosses the proportion of the compared of the compared of the proportion of the compared of the proportion of the compared of the com

References: 1. Data on file. Pizer Inc., New York, NY. 2. Simpson GM, Glick ID, Weiden PJ, Bromano SJ, Siu CO. Randomized, controlled, double-blind multicenter comparison of the efficacy and tolerability of ziprasidone and olanzapine in acutely ill inpatients with schizophrenia or schizoaffective disorder. Am J Psychiatry, 2004;161:1837-1847. 3. Addington DEN, Pantells 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 trail. J Clin Psychiatry, 2004;65:1624-1633. 4. Simpson GM, Weiden P, Pigott T, Worray 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.

Treat schizophrenia with the body in mind

COMPARABLE EFFICACY

Consistent results in head-to-head studies1-3

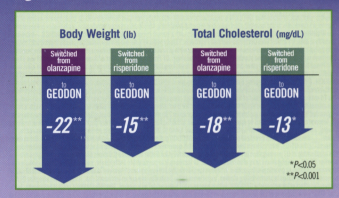


A 6-week, double-blind, randomized study of GEODON vs olanzapine and an 8-week, double-blind, randomized study of GEODON vs risperidone.

- 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

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}



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 ≥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%).

Please see brief summary of prescribing information on adjacent page.