## CALL FOR PAPERS

CNS Spectrums is accepting submissions of case reports, review articles, and original research on a variety of neuroscientific and clinical neuropsychiatric topics.

#### **Examples of topics include:**

- Clinical interface of psychiatry and neurology
- Neurology and neuropsychiatry in a clinical setting addressing spectrum disorders
- Applications of psychopharmacology and pharmacokinetics across the neuropsychiatric spectrum

Especially encouraged are papers covering comorbidities in neurologic disorders (eg, epilepsy with schizophrenia). Other crossover manuscripts geared to deepening the clinician's understanding of neuropsychiatric disorders and treatments will be given highest priority. (Please see the Author Guidelines at www.cnsspectrums.com/aspx/ authorquidelines.aspx).

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CNS Spectrums has the largest circulation among Index Medicus-approved publications with a monthly readership of 50,000 neurologists and psychiatrists worldwide.

Submissions should be sent to Eric Hollander, MD, Editor (In Europe, to Joseph Zohar, MD, International Editor), c/o Virginia Jackson, Acquisitions Editor, CNS Spectrums, c/o MBL Communications, 333 Hudson Street, 7th Floor, New York, NY 10013, E-mail: vi@mblcommunications.com.





A Global Commitment to Advancing CNS Science, Clinical Practice, and Evidence-Based Medicine

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 to between 1.5 to 1.7 times that seen in placebo-treated patients. Ower the course of a rylical 10 week controlled trial. In relate of death in drug-treated patients. Ower the course of a rylical 10 week controlled trial. In relate 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 actients with Dementia-Related Psychosis 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 teatures. GEODON® (ziprasidone mesylate) for injection is indicated for acute agitation in schizophrenic patients.

SCINZAPINENTED BANGINGS.

CONTRAINDICATIONS — OT Prolongation: Because of GEODON'S dose-related prolongation of the QT interval and the known association CUN HANDLCA IUNS — 11 Prolongation: because of sEUDON's dose-related prolongation or the U1 interval and the known association of fatal arrivythinias with OT prolongation by some other drugs, GEODON is contraindicated in patients with a known bistory of OT prolongation (including congenital long OT syndrome), with recent acute myocardial infarction, or with uncompensated heart failure (see WARNINGS). Pharmacokinetic/pharmacodynamic studies between 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-arrivythmics, mesoridazine, thioridazine, chlorpromazine, droperidol, be given with doteflide, sotabl, quinidine, other Class is and III anti-arritythmics, mesondazine, chioridazine, chioripromazine, droperidol, pimozode, sparfloxacin, gatificiazion, mostificazion, halofantrine, melloquine, pentamidine, arsenic trioxide, levomethady acetale, dolasetron mesylate, protucol, or tarcrilimus. GEODON is also contraindicated that have demonstrated QT profuncian as one of their pharmacodynamic effects and have this effect described in the full prescribing information as contraindication or a boxed or bolded warning (see WARNINGS). GEDOON is contraindicated in individuals 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 drugs their antipsychotic drugs are at an increased risk of death compared to placebo. GEODON (dyrasidone) is not approved for the treatment of patients with dementia-related psychosis (see Boxed Warning). AT Prolongation and Risk of Sudden Death: GEODON use should be avoided in combination with other drugs that are known to protong the QT, interval. Additionally, clinicians should be aefer to the identification of other drugs that have been consistently observed to protong the QT, interval. Such drugs should not prescribed with GEODON. A study directly comparing the QT/QT-prolonging effect of GEODON with several other drugs effective in the treatment of schizophrenia was conducted in patient volunteers. The mean increase in QT, from baseline for GEODON ranged from approximately 9 to 14 mase greater than for four of the comparator drugs (risperidone, olanzapine, quetlapine, and hatoperidol), but was approximately 14 mase class than the prolongation observed for thioridazine. In his study, the effect of GEODON on QT, length was not augmented by the presence of a metabolic inhibitor (teleconazione 200 mg bild). In placebo-confolied trials, Geolon increased the clinically relevant threshold of 500 msec. In the GEODON patients, neither ca electrocardiograms of 2/2988 (0.06%) GEODON patients and 1/440 (0.23%) placebo patients revealed OT, intervals exceeding the potentially clinically relevant threshold of 500 msec. In the GEODON patients, neither case suggested and led GEODON. Some drugs that prolong the OT/OT, prolongation to torsade de pointes is clearest for larger increases (20 msec and greater) but it is possible that smaller QT/OT, prolongations may also increase risk, or increase it in susceptible individuals, such as those with hypokalemia, hypomagnesemia, or genetic precisposition. Although torsade de pointes has not been observed in association with the use of GEODON at recommended doses in permarketing studies, experience is too limited to rule out an increased risk. A study existing the QT/OT, prolongalions may also increase risk, or increase it in susceptible individuals, such as those with hypokalemia, hypomagnesemia, or genetic precisposition. Although torsade de pointes has not been observed in association with the use of GEODON at a recommended doses in permarketing studies, experience is too limited to rule out an increased risk. A study experiment to the time of the commended threapen obtained at the time of maximum plasma concentration following the violence of GEODON (20 mg then 30 mg) or hatoperidol (7.5 mg then 10 mg) given four hours a part. Note that a 3 mg dose of intramuscular GEODON is 50% higher than the recommended threapenite dose. The mean change in QT, from baseline value using a sample-based correction that removes the effect of heart rate on the QT interval. The mean increase in QT, from baseline to GEODON was 4.6 msec following the first injection and 14.7 msec following the second injection. In this study, no patient had a QT, interval exceeding 500 msec. As with other antipsychotic drugs and placebo, sudden unexplained deaths have been reported in patients taking GEODON at a considered doses. The premarketing experience for GEODON did not reveal as nexcess of mortality for GEODON Compared to other antipsycho since recurrences of wins have been reported. Internet proximental (10), A syndrome or potentially interesting, involuntary, cystam movements may develop in patients undergoing treatment with antipsycholic drugs. Although the prevalence of TD appears to be highest among the elderly, especially elderly women, it is impossible to rely upon prevalence estimates to predict, at the inception of antipsycholic 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. ##pperpicemia and Diabetes Melitius: Hyperglycemia-related adverse events, sometimes serious, have been reported in patients treated with applical and hipsychotics. 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. \*Patients\* ITT in premarketing traits, about 5% of GEODON patients developed rash and/or urlicaria, 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 litness, e.g., elevated WBCs. Most patients improved promptly upon treatment with antihistamines or steroids and/or upon discontinuation of GEODON, and all patients were reported to recover completely. Upon appearance of rash for which an alternative eliopsy cannot be identified. GEODON should be discontinued. Orthostatic Hypotension: GEODON may induce orthostatic hypotension associated with dizziness. Lachycardia, and in some patients, syncope, sepsecially during the initial dose-thristion period, problems fredering its c<sub>17</sub> 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 (instory of myocardial rediscose patients to hypotension (dehvidration, hypovolemia, and patients to the production (dehvidration, hypovolemia, and patients of the production (dehvidration) in the patients of the production (dehvidration). 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, hypovoienia, and treatment with antihyportensive 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 antipoychotic drugs, GEODON should be used cautiously in patients with a history of seizures or with conditions that potentially lover 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. <a href="Dysphagic Esophageal dysmotility">Dysphagic Esophageal dysmotility</a> and spiration 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, and GEODON 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). <a href="Hyperprojectimenias">Hyperprojectimenias</a>, As with other drugs that antagonize dopamine Dy receptors, GEODON elevates productin levels in humans. Tissue culture experiments indicate that approximately on third of human brasa 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. Neither clinical studies nor epidemiologic studies conducted to date have shown an association between chronic administration of this class of drugs and tumoriquensis in humans. Tesualisable evidence is considered too limited to be conclusive at this time. Potential for Cognitive Invitro, a factor of potential importance if the prescription of these drugs is contemplated in a patient with previously detected breast cancer. Nether clinical studies nor epidedine, locations, conducted to date have shown an association between chronic administration of this class of drugs and tumorigeness in humans; the available evidence is considered too limited to be conclusive at this time. <u>Potential for Cognitive</u> and <u>Motor Impairment.</u> Somnolence was reported in 14% of 6EODON patients vs. 7% of placebo patients. In the 4-and 6-week placebo-controlled trails, somnolence was reported in 14% of 6EODON patients vs. 7% of placebo patients. Somnolence led to discontinuation in 0.3% of patients in short-term clinical trails. Since 6EODON 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 verbile (including automobiles) or operating hazardous machinery until they are reasonably certain that 6EODON therapy does not affect them adversely. <u>Prapism.</u> One case of priapism was reported in the premarkening database. <u>Body Temperature Regulation.</u> Although not reported with GEODON in premarketing raises through the prescriptions should be written for the smallest clausified or supervision of high-risk patients which certain concomitant systemic illness so hould be written for the smallest claunity of capacities consistent with good patient management to relative to proportion and orthostatic hypotension with 6EODON in patients with ocratic concomitant systemic illnesses is limited. GEODON was not been evaluated or used to any appreciable extert in patients with ocratic chapters with the certain concomitant systemic illnesses of the risk of 10 fp. prolongation and orthostatic hypotension with 6EODON in patients with ocratic comparison of intramuscular in hypotension in PRECAUTIONS). *Information for Patients*: To ensure safe and effective use of 6EDOON, he nonly symptoms reported were minimal

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 plotassium and magnesium should be repleted before treatment. Patients who are started on disturbing GEODON therapy need periodic monitoring of serum potassium and magnesium. Discontinue GEODON in patients who are found to have persistent OT, measurements. SOO mass (see WaRAINGS). *Drug Internactions*: (1) GEODON should not be used with any drug that prolongs the OT internal. (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 antihyportensive agents. (4) GEODON may antagonize the effects of levodops and dopamine agonists. <u>Effect of Other Drugs on GEODON</u>, *Cartamazepine*. 200 mg bid for 21 days, resulted in a decrease of approximately 35% in the AUC of GEODON. *Rebocorazole*, a potent inhibitor of CYP344. 400 mg of for 5 days, increased the AUC and Cm<sub>max</sub> of GEODON by about as 5%. 40%. *Circumbine*, 800 mg of for 2 days, did not 2 days. did not affect GEODON pharmacokinetics. Population pharmacokinetic analysis of sorbizophering patients in controlled chiral atrials has not revealed any clinically significant pharmacokinetic interactions bear propriations of drugs cleared primarily by CYP1A2, CYP2OB, CYP2OB, and CYP3A4, and little potential for drug interactions with GEODON do mgl and decomposed to the displacement. GEODON 400 mgl bid did not affect the pharmacokinetics of concomitantly administered or ordrazoptive. 2 study in normal bacter the stagory-state level or renal clearance of lithium to GEODON 20 mg bid did not affect the pharmacokinetics of concomitantly administered ordrazoptives. 2 study is normal bacter to the stagory-state level or renal clearance of lithium composition. information and instructions in the Patient Information Sectionshould be discussed with patients. Laboratory Tests: Patients being considered ESDON due to displacement. ESDON 40 mg bid administered concomitantly with ithium-450 mg bid for 7 days did not affect the steady-state level or renal clearance of lithium. ESDON 20 mg bid did not affect the pharmacokinetics of concomitantly administered or acontracephives, ethiny estradiol (0.03 mg) and levonorgestrel (0.15 mg). Consistent with in vitro results, as tudy in mormal healthy volunteers showed that ESDON did not after the metabolism of devironethorphan, a CYP2D6 model substrate, to its major metabolite, devironethorphan devertorphan ratio. Carcinogenia healthy volunteers showed that GEODON did not after the metabolism of the uniany devironethorphan/dextrorphan ratio. Carcinogenia maintenance of the unitary devironethorphan/dextrorphan ratio. Carcinogeniasis, Mutagenesis, Impairment of Fartility. 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 re-related increases in serum prolactin were observed in a 1-month dietary study in female, but not male, mice. ESCODON had no effect on serum prolactin in ratis in a 5-week dietary study at the doess that were used in the carcinogenicity study. The relevance for human risk of the findings of prolactin-mediated endocrine humors in rodents is unknown (see Hyperprolactinemia). Mutagenesis; There was a reproducible mutagenic response in the American sassay in human hymphocytes. Impairment of Fertility. GEODON increased time to copulation in Sprague-Dawley rats in Not entrility and early and the control of metabolic activation. Positive results were obtained in both the in Mgg/day (25 to 8 times the MRHD or 200 mg/day or a mg/m² basis). There was no effect on fertility af dompkyday (25 times the MRHD on a mg/m² basis). The relative to control the male relative service of the potential benefit justifies the potential risk to the fetus. Labor and Delivery. The effect of ECDON in clabor and delivery in humans is unknown. Musing Traks for schizoprima (a) pool of two 5-week (nath two 4-week fixed-does this) and bipolar mania (a pool of two 3-week fixed for obsertials) in which 6EDON was administered in doses ranging from 10 to 200 mg/day. Adverse Events Associated with Discontinuation: Schizoprima: Approximately 4.15 (2070z) of 1000 th-reated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 2.2% (6073) 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 PEGEAUTIONS). Bipolar Mania: Approximately 6.5% (18279) of EEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 2.2% (6073) on placebo. The most common events associated with dropout was rash, adverse event compared with about 2.7% (6736) on placebo. The most common events associated with group of the term of the placebo patient schi or dystonia and rash of the placebo patients feet propout in the EEDOOM-treated patients were alkathisia, anxiety, depression, dizziness, dystonia, rash and vorniting, with 2 dropouts for each of these events associated with GEODON in compared to one placebo patient schi or dystonia and rash (1%) and no placebo patients for the remaining adverse events. Adverse Events at an incidence 25% and at Least Twice the Rate of Placebo: The most commonly observed adverse events associated with GEODON in bippar mania thials were somnolence (31%), attention of the compared to the compared to the compared to the compared to the compared adverse events that occurred during acute therapy, including only those events that occurred during acute therapy, including only those events that occurred during acute therapy including any those events that occurred during a compared to the compared to t hypoglycemia, hypomatremia, hypoproteinemia, glucose tolerance decreased, gout, hyperchloremia, hyporuticemia, hypoproteinemia, hypoproteinemia, educose tolerance decreased, gout, hyperchloremia, hyporuticemia, hypoproteinemia, Incidence 3-1% in Short-Term Fixed-Dose Inframuscular Trials: The following list enumerates the treatment-emergent adverse events that occurred in a 1% of 6EODON patents (in the higher dose groups) and at least twice that of the lowest inframuscular planet is not higher dose groups) and at least twice that of the lowest inframuscular planet in the properties of the prop

Revised May 2005

### Control acute agitation with

# GEODON® for Injection | ziprasidone mesylate|

In schizophrenia...

# Rapid improvement with low EPS<sup>1,2</sup>

- Significant control achieved between 15 and 30 minutes\* after injection<sup>1,3</sup>
- Proven advantages over haloperidol IM
  - twice the improvement as measured on the BPRS $^{4\dagger}$
  - significantly lower incidence of movement disorders<sup>2‡</sup>
- Smooth transition, with continued improvement, from IM to oral therapy<sup>2,4</sup>
- 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<sub>C</sub> 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.