3rd INTERNATIONAL CONGRESS ON

W MEN'S MENTAL HEALTH MELBOURNE AUSTRALIA

17-20 March 2008

Opening Minds to Women's Mental Health

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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).

MBL Communications, Inc., is proud of the 2005 ISI Journal Citation Reports' impact factor for CNS Spectrums. The current impact factor is 2.037 for CNS Spectrums and is based on a total of 580 citations.

CNS Spectrums has the largest circulation in the nation among peer-reviewed, Index Medicus-approved neuropsychiatric 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: vj@mblcommunications.com or submitted electronically at www.cnsspectrums.com.

PRIMARY PSYCHIATRY CNS SPECTRUMS Psychiatry Weekly.

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

Scope of Manuscripts

CNS Spectrums will consider and encourages the following types of articles for publication:

- 1. <u>Original Research</u> presents methodologically sound original data.
- <u>Reviews</u> are *comprehensive* articles summarizing and synthesizing the literature on various neuropsychiatric topics and presented in a scholarly and clinically relevant fashion. Diagnostic and treatment algorithms should be designed to aid the clinician in diagnosis and treatment.
- 3. <u>Case Reports</u>, single or multiple, are encouraged for publication.
- Letters to the Editor will be considered and are encouraged for publication. All letters will be edited for style, clarity, and length.

Manuscript Submission

General Information Two copies of the manuscript with a letter on the author's letterhead should be submitted to **Eric Hollander**, **MD**, **Editor** (or, in Europe, to Joseph Zohar, MD, International Editor), c/o MBL Communications, 333 Hudson Street, 7th Floor, New York, NY 10013. Authors are also required to submit their manuscripts on computer disk in Microsoft Word format. Disks should be labeled with the word processing program, title of paper, and lead author's name. Accepted manuscripts will be edited for clarity and style.

eSubmissions Now Available Please go to http://cnsspectrums.com/aspx/authorguidelines.aspx. E-mail your your manuscript to Virginia Jackson, Acquisitions Editor, *CNS Spectrums*, at vj@mblcommunications.com.

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Peer Review Authors must provide five to 10 names of qualified potential reviewers with no conflict of interest in reviewing the work. Contact information with affiliations and e-mail address should be included. Peer review is anonymous.

Manuscript Preparation

Length Reviews and Original Research should not exceed 5,000 words (excluding References). Diagnostic and treatment algorithms should contain an introduction, flowcharts or a series of graphs, and a concise summary. At least 2 tables or figures are required. Letters should not exceed 1,500 words. Single-Case Reports should not exceed 3,750 words and may be submitted with a photograph, if applicable.

Please note: If your article is Original Research, it should be formatted as: Abstract; Introduction, Methods; Results; Discussion; Conclusion; References (numbered and comprehensive list).

Spacing and Pagination Manuscripts should be doublespaced and numbered.

Abstract Authors must provide a brief abstract of 100–200 words.

Focus Points Please provide 3–6 focus points that begin with an action verb and specify what the reader should know after reading the article.

Learning Objectives Authors are required provide 3–6 learning objectives, which begin with an action verb and specify what the reader should know after reading the article. See the following examples:

Upon the completion of this lecture the participants will be able to:

- List four causes of aplastic anemia
- Give an example of the effect of a strong alkali reacting with human tissue
- Calculate the amount of AIV fluid necessary to replenish a dehydrated patient

Needs Assessment Please provide a brief summary (35–50 words) outlining the educational needs and reasons for reading the article. It should address a deficit or gap in knowledge, skills, attitudes, and/or behavior among the expected readers about the main topic of the article. It should justify the reasons for focusing on the given topic and offering it as a CME activity. Reasons would include recurrent discussions with colleagues about the topic, new therapy or treatment techniques, new data published, "hot topic" in the field, clinical trials in progress, etc.

Figures/Tables Please provide at least 2 original figures and/or tables.

References Please use American Medical Association style. References should be superscripted in text, then numbered, and comprehensive in list. For example:

- 1. Jones J. Necrotizing Candida esophagitis. JAMA. 1980;244:2190-2191.
- Stryer L. Biochemistry. 2nd ed. San Francisco, Calif: WH Freeman Co; 1980:559-596.
- Alzheimer's Disease Cooperative Study. Valproate protocal. Available at: http://adcs.ucsd.edu/VP_Protocol.htm. Accessed October 15, 2003.

Continuing Medical Education Authors must submit 6 multiple-choice questions with answers.

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Disclosure of Commercial and Non-Commercial Interests

Authors must include a statement about all forms of support, including grant and pharmaceutical support, affiliations, and honoraria received for past and present material. Such information may, at the editor's discretion, be shared with reviewers. If the article is accepted for publication, the editors will consult with the authors as to whether this information should be included in the published paper.

Submission Checklist

- Original manuscript plus one copy, with cover letter on author's letterhead
- Copies of permission letters to reproduce previously published and unpublished material
- A brief abstract of the article
- □ Six multiple-choice CME questions with answers
- 3-6 focus points that dictate the main focus of the manuscript in bulleted format
- 3–6 learning objectives, which begin with an action verb and specify what the reader should know after reading the article
- Disk labeled with the word-processing program, title of paper, and lead author's name
- Names and affiliations of 3–5 potential peer reviewers

2008 INTERNATIONAL ANXIETY DISORDERS SYMPOSIUM

March 17–18, 2008 at the Arabella Sheraton Hotel, Cape Town, South Africa*

A Taste of the Programme:

Chairperson: Dan J. Stein, MD, PhD

• Anxiety Disorders in Schizophrenia David Castle, MD University of Melbourne (Australia)

• Endophenotypes in Obsessive-Compulsive Disorder Naomi Fineberg, MD University of Hertfordshire (United Kingdom)

- Substance Abuse and Anxiety Disorders Marc Schuckit, MD University of California at San Diego (United States)
- Optimising Diagnosis in the Community Christer Allgulander, MD Karolinska Institute (Sweden)

• Repetitive Transcranial Magnetic Stimulation in Anxiety Disorders Jack van Honk, PhD Utrecht University (Netherlands)

And a range of other expert speakers

Dates to Remember:

- November 9, 2007 Closing date for electronic abstracts
 - January 11, 2008 Closing date for early registration
- February 22, 2008 Closing date for symposium registration

For more information and to register for the conference, please visit: www.mentalhealthsa.co.za/anxietyconference/registration.php or contact: <u>Arlene Kleinhans at arlene@sun.ac.za</u>

* <u>Please note this conference takes place immediately after the International Society for</u> <u>Affective Disorders meeting taking place March 14–17 at the Arabella Sheraton Hotel</u>

CNS Spectr 12:9

BRIEF SUMMARY. See package insert for full prescribing information

Increased Montally in Edistry Patients with Dementia-Related Psychosis. Elderly patients with dementia-related psychosis treated with atypical antipsychotic damps are at an increased risk of death compared to placebe. Analyses of seventies n lacebe controlled trials (model 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 placebot-headed patients. Over the course of a typical 10 week controlled trial, the rate of death in drug-treated patients are weeks and the second of the second patients of between 1.6 to 1.7 times that seen 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., head failure, sudden death) or intectious (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

The sense cardworkscale (e.g., feet nature, souden beauty of microlog (e.g., preunothal) in fault. BCDOM (dpressione) is not approved in the treatment of schizophrenia and acute marker or mixed episodes associated with biplar disaders with a vertical schizophrenia and acute marker or mixed episodes associated with biplar disaders with a vertical schizophrenia patients. COODM* (dpressione mesylate) for injection is indicated for acute agitation in schizophrenia patients. COODM* (dpressione mesylate) for the convertige agitation in schizophrenic patients. COOTMANDCATIONS— of Prolongation by some other drugs, GEODON's does related prolongation of the CT interval and the known association of tatal arriythmias with OT prolongation (and OT Syndrome), with resent acute mexorcatic line tracture, GEODON should not be own with dotellide, schizophrenic patients. COODN and other drugs that prolong the CT interval acute be colded. Therefore, GEODON should not be own with dotellide, schizophrenic agitate prove that acute (adia acute agitation) in produces and the convertigent and the schizophrenic patients. GEODON is also contraindicated with drugs tath ave demonstrated OT prolongation as ore of their pharmacolymanic effects and have been constraindicated with a low down hypersansitivity to the product. WARNINGS - MERCENDN is also contraindicated with a low down hypersansitivity to the product. WARNINGS - MERCENDN is a schizophrenic s potentially fala symptom complex sometimes referred to as Neuroleptic Maignant Syndrome (MMS) 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 drugs and there drugs not essential to encurrent therapy. (2) intensive symptomatic treatment and medical monitoring; and (3) treatment and 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. **Tartine Dyskinesia (TD)**: A syndrome of potentially irreversite, involutary, dyskinetic movements may develop in patients undergoing treatment with antipsychotic drugs and the series 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 TD. If signs and symptoms of TD appear in a patient on GEDODM, drug discontinuation should be considered. *Hyperglycemia and Diabetes Mellitus*: Hyperglycemia-retaid adverse events, sometimes serious, have been reported in and it is not known if GEDODM, is associated with these events. Faiteris treated with antipsychotic should be monitored for symptoms of hyperglycemia. **PRECAUTIONS**— *General*: Bash; In premarketing trials, about 5% of GEDODM, patients developed rash and/or unicrait, which generalized and patients were reported to recover completely. Upon appearance of rash was dose reliated, although the finding might also be explained by longer exposure in higher-dose patients. Several patients with rash had signs and symptoms of associated with diziness, e.g., elevated WBCS. Most patien trastment with anthropertures enderations), Siggings, In clinical trials, secures courred in 0.4% of ECDON patients. There with a solution trians of these sees. As with other anticocyclic days, ECDON should be anticocyclic days, and sprator have been associated with analyse/botic days sea. As with other anticocyclic days, ECDON should be anticocyclic days, and sprator have been associated with analyse/botic days sea. As with other anticocyclic days, and sprator have been associated with analyse/botic days sea. As with other anticocyclic days, and sprator have been associated with analyse/botic days sea. As with other anticocyclic days as as board WhANNE, WANNES, MANNES, WANNES, MANNES, MANNES,

<text> 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

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

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

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

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



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

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

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

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

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

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

Please see brief summary of prescribing information on adjacent page.