AUTHOR GUIDELINES

Introduction

CNS Spectrums is an Index Medicus journal that publishes original scientific literature and reviews on a wide variety of neuroscientific topics of interest to the clinician. CNS Spectrums will publish 12 issues in 2003. As the immense prevalence of comorbid diseases among patients seen by psychiatrists and neurologists increases, these physicians will jointly diagnose and treat the neuropsychiatrically ill. Our mission is to provide these physicians with an editorial package that will enhance and increase their understanding of neuropsychiatry; therefore, manuscripts that address crossover issues germane to neurology and psychiatry will be given immediate priority.

Scope of Manuscripts

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

Original Reports: Original reports present methodologically sound original data.

Reviews: Reviews are overview articles that summarize and synthesize the literature on various topics in a scholarly and clinically relevant fashion. Suitable topics include mood disorders, schizophrenia and related disorders, personality disorders, substance-use disorders, anxiety disorders, neuroscience, psychosocial aspects of psychiatry, child psychiatry, geriatric psychiatry, and other topics of interest to clinicians. Original flowcharts designed to aid the clinician in diagnosis and treatment will be considered for publication in reviews and are encouraged.

Case Reports: Single or multiple case reports will be considered for publication.

Letters to the Editor: Letters will be considered for publication.

Manuscript Submission

General information: Two copies of the manuscript with a letter on the author's letterhead should be submitted to Jack M. Gorman, Editor (or, in Europe, to Joseph Zohar, International Editor), c/o MBL Communications, 333 Hudson Street, 7th Floor, New York, NY 10013; (F) 212.328.0600. 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 and letters will be edited for clarity and style.

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Peer review: Authors must provide five names of parti-cularly qualified potential reviewers with no conflict of interest in reviewing the work. Contact information, including complete address, phone, fax numbers, E-mail address, and affiliations, should be included. The corresponding author will be notified by the editors when a decision regarding acceptance has been made. Peer review is anonymous.

Manuscript Preparation

Length: Reviews and Original Reports should not exceed 5,000 words (excluding References). 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. Diagnostic/treatment algorithms (see Reviews) should contain an extensive introduction, flowchart or series of graphs that fill 8–12 journal pages, and a concise summary.

Spacing: One space should be left after commas and periods. Manuscripts should be double-spaced.

Abstract: Authors must provide a brief abstract.

References: American Medical Association style. See the following examples:

- 1. Jones J. Necrotizing Candida esophagitis. JAMA. 1980;244:2190-2191.
- 2. Stryer L. Biochemistry. 2nd ed. San Francisco, Calif: WH Freeman Co; 1980:559-596.

Continuing Medical Education: Authors must submit four multiple-choice questions (two Type A and two Type K), with answers.

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Disclosure of Commercial Interests

Authors must include a statement about all forms of support, including grant and drug company support. 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
- ☐ Four CME multiple-choice questions with answers
- ☐ Disk labeled with the word processing program, title of paper, and lead author's name
- ☐ Names and addresses of five potential reviewers

GUIDE TO DSM-IV AND ICD-10 CODES

Describe of the Make Inner Too With Early One of With Described	DSM-IV	ICD-10
Dementia of the Alzheimer Type, With Early Onset With Depressed Mood Specify if: With Behavioral Disturbance	290.13	F00.03
Dementia of the Alzheimer's Type, With Late Onset With Depressed Mood Specify if: With Behavioral Disturbance	290.21	F00.13
Delirium Due to: Indicate General Medical Condition	293.0	F05.0
Psychotic Disorder Due to: Indicate General Medical Condition With Delusions With Hallucinations	293.81 293.82	F06.2 F06.0
Mood Disorder Due to: Indicate General Medical Condition	293.83	F06
Anxiety Disorder Due to: Indicate General Medical Condition Amnestic Disorder Due to: Indicate General Medical Condition	293.89 294.0	F06.4 F02.8
Dementia NOS	294.8	F03
Amnestic Disorder NOS	294.8	R41.3
Schizophrenia—Disorganized Type	295 295.10	F20 F20.1
Schizophrenia—Catatonic Type	295.20	F20.2
Schizophrenia—Paranoid Type Schizophrenia—Residual Type	295.30 295.60	F20.0 F20.5
Schizoaffective Disorder	295.70	F25
Schizophrenia—Undifferentiated Type Major Depressive Disorder	295.90 296	F20.3 F32
Bipolar I Disorder	296	F30
Bipolar Disorder NOS Bipolar II Disorder	296.80	F39
Bipolar II Disorder Mood Disorder NOS	296.89 296.90	F31.8 F39
Psychotic Disorder NOS	298.9	F29
Autistic Disorder Asperger's Disorder	299.00 299.80	F84 F84.5
Pervasive Developmental Disorder NOS	299.80	F84.9
Anxiety Disorder NOS	300.00	F41.9 F41
Panic Disorder Without Agoraphobia Generalized Anxiety Disorder	300.01 300.02	F41.1
Dissociative Identity Disorder	300.14	F44.81
Dissociative Disorder NOS Factitious Disorder NOS	300.15 300.19	F44.9 F68.1
Panic Disorder With Agoraphobia	300.21	F40.01
Agoraphobia Without History of Panic Disorder Social Phobia	300.22 300.23	F40 F40.1
Specific Phobia	300.29	F40.1 F40.2
Obsessive-Compulsive Disorder	300.3	F42.8
Dysthymic Disorder Depersonalization Disorder	300.4 300.6	F34.1 F48.1
Body Dysmorphic Disorder	300.7	F45.2
Somatization Disorder Somatoform Disorder NOS	300.81 300.81	F45. F45.9
Cyclothymic Disorder	301.13	F34
Alcohol Dependence Cocaine Dependence	303.90 304.20	F10.2 F14.2
Cannabis Dependence	304.20	F14.2 F12.2
Amphetamine Dependence	304.40	F15.2
Alcohol Abuse Cannabis Abuse	305.00 305.20	F10.1 F12.1
Cocaine Abuse	305.60	F14.1
Amphetamine Abuse Stuttering	305.70 307.0	F15.1 F98.5
Anorexia Nervosa	307.0	F50
Tic Disorder NOS	307.20	F95.9
Tourette Disorder Primary Insomnia	307.23 307.42	F95.2 F51.0
Primary Hypersomnia	307.44	F51.1
Sleepwalking Disorder Dyssomnia NOS	307.46 307.47	F51.3 F51.9
Nightmare Disorder	307.47	F51.5
Parasomnia NOS Eating Disorder NOS	<u>307.47</u> 307.50	F51.8 F50.9
Bulimia Nervosa	307.50	F50.2
Feeding Disorders of Infancy or Early Childhood	307.59	F98.2
Communication Disorder NOS Posttraumatic Stress Disorder	307.9 309.81	F80.9 F43.1
Depressive Disorder NOS	311	F32.9
Impulse-Control Disorder NOS Pathological Gambling	312.30 312.31	F63.9 F63.0
Pyromania	312.33	F63.1
Kleptomania Trichotillomania	312.34 312.39	F63.2
Disruptive Behavior Disorder NOS	312.39 312.9	F63.3 F91.9
Attention-Deficit/Hyperactivity Disorder, Combined Type	314.01	F90
Attention-Deficit/Hyperactivity Disorder NOS Learning Disorder NOS	314.9 315.9	<u>F90.9</u> F81.9
Developmental Coordination Disorder	315.4	F82
Narcolepsy Sleep Disorder Due to: Indicate General Medical Condition	347 780	G47.4 G47
Delirium NOS	780.09	F05.9

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1 2 3 4 5 CME	4. On a scale of 1 to 5 (1=Incomplete, 5=Comprehensive), how would you describe the depth of coverage for this issue?
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The Neurology of Behavior	1 2 3 4 5
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INUICATIONS AND USAGE
SEROULE Is indicated for the treatment of schizophrenia.
The efficacy of SEROULE in schizophrenia was established in short-term (6week) controlled trials of schizophrenic inpatents (See CLINICAL PHARMACOLOGY).
The effectiveness of SEROULE in long-term use, that is, for more than 6 weeks,
as not been systematically evaluated in controlled trials. Therefore, the physician
who elects to use SEROULE for extended periods should periodically re-evaluate
the long-term selfuness of the drug for the individual patient
CONTRANDICATIONS
SEROULE Is contrainficated in individual patient.

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Neurolepic Mailgnand Syndrome: (MMS) A potentially fatal symptom complex comertimes referred to as Neurolepic Mailgnand Syndrome (MMS) local seem resported with the process of the common complex comertimes referred to all and that with SERODUEL. Clinical manifestations of IMMS are hyperpyrexa, muscle rigidity, altered mental status, and evidence of automolic instability regular professor by donor processor. In the dispension of the processor of the dispension of the syndrome and therefore the dispension of the syndrome and therefore the dispension of

SEROQUEL® (quetiapine fumarate) Tablets

have been associated with antipopychoted ruppus. Apparation presumons is a common cause of metodelly and controlly on addrey proteins. In particular tools with motivation of the controlly of th

SEROUGE: We present the formative of tables.

Natising Mothers: SEROUEL: was excreted in milk of treated animals during lactation its not known if SEROUEL is excreted in human milk. Its recommended that women receiving SEROUEL is pediatric patients have not been established. Berlatic Use: Of the approximately Adol patients in clinical studies with SEROUEL. 8% reglowanted by Adol patients in clinical studies with SEROUEL. 8% reglowanted by Adol patients in clinical studies with SEROUEL. 8% reglowanted by Adol patients in clinical studies with SEROUEL. 8% region there was no indication of any different iderability of SEROUEL in the sidenty compared to younger adults. Nevertheless, the presence of factors that might decrease paramacokinetic clearance, increase the pharmacokynamic response to SEROUEL or cause poorer tolerance or or orbustass; should lead to consideration of a lower starting dose, slower titration, and careful monitoring during the initial dosing period in the eddery. The mean plasma clearance of SEROUEL was reduced by 30% to 50% in elderly patients when compared to younger patients. younger patients.
ADVERSE REACTIONS

ADVERSE REACTIONS
Adverse Events Decurring at an Incidence of 1% or More Among SEROQUEL
Treated Patients in Short-Term, Placebo-Controlled Trials: The most commonly
observed adverse events associated with the use of SEROQUEL (incidence of 5% or
greater) and observed at a rate on SEROQUEL at least twee that of placebo were
dizziness (10%), postural hypotension (7%), dry mouth (7%), and dyspepsia (6%).
The following trathent-emergend adverse experiences occurred at an incidence rate
of 1% or more, and were at least as frequent among SEROQUEL treated patients,
treated at dossor 57 5 mg/day or greater than among placebo treated patients in
3- to 6-week placebo-controlled trials.

Body as a Whotel Headach A Athenia Abdoniral pain Bod Acid Four March

Body as a White-Headache, Ashina, Abdominal pain, Back pain, Fever Nerveus System: Somnolence, Dizziness; Digestive System: Constipation, Dry Mouth, Dysopejac, Gardiovascular System: Postural typotenson, Tachycardio, Metabolic and Mutritional Disorders: Weight pain; Staff and Appendagues: Hash, Respiratory System: Rininis; Speala Sanses: Ear pain Feveris for which the SEROULE! incidence was equal to or less than placebo and staff and table but incidence was equal to or less than placebo and staff and table but incidence was equal to or less than placebo and staff and table but incidence was equal to or less than placebo and staff and the stable but incidence was equal to or less than placebo and staff and the stable but incidence was equal to or less than placebo and staff and the stable was expendenced and the staff and sta

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The most common adverse events associated with the use of SEROQUEL are dizziness (10%), postural hypotension (7%), dry mouth (7%), and dyspepsia (6%). The majority of adverse events are mild or moderate.

In 3- to 6-week, placebo-controlled trials, the incidence of somnolence was 18% with SEROQUEL vs 11% with placebo.

As with all antipsychotic medications, prescribing should be consistent with the need to minimize the risk of tardive dyskinesia, seizures, and orthostatic hypotension.

References: 1. Prescribing Information for SEROQUEL® (quetiapine furnarate), Rev 1/01, AstraZeneca Pharmaceuticals LP, Wilmington, Delaware. 2. Data on file, IMS data, AstraZeneca Pharmaceuticals LP, Wilmington, Delaware.



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