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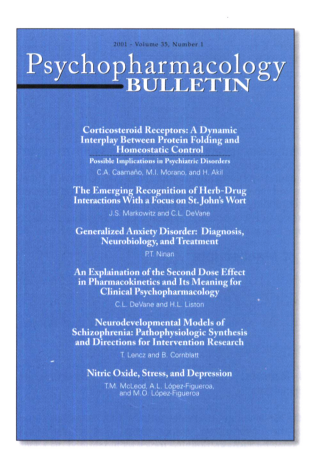
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Introduction

C S S Þ е s is a peer-reviewed journal с publishes original scientific literature and reviews on a wide variety of neuroscientific topics of interest to the clinician. C S е m s publishes 12 issues in 2001. prevalence As the immense 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

C N S p p e c t r u m s 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 Submissions

General information: Four copies of the manuscript should be submitted to Jack M. Gorman, editor (or, in Europe, to Joseph Zohar, international editor), c/o MedWorks Media, 333 Hudson Street, 7th Floor, New York, NY 10013; T: 212.328.0800, F: 212.328.0600. Authors are required to submit their manuscripts on computer disks. If possible, please provide them in MS Word for Windows in either a Macintosh or IBM format. (Saving the file in a lower version, eg, MS Word 3.0, is also encouraged.) Disks should be labeled with the word-processing program, title of paper, and first author's name.

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Peer review: Authors should provide five names of particularly 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. Accepted manuscripts and letters will be edited for clarity and style.

Manuscript Preparation

Length: Reviews should not exceed 20 manuscript pages (10,000 words). Original reports should not exceed 15–25 manuscript pages (6,250 words, maximum). Letters should not exceed 2–6 manuscript pages (1,500 words, maximum). Single case reports should not exceed 10–15 manuscript pages (3,750 words, maximum) and may be submitted with a photograph, if applicable. Diagnostic/treatment algorithms (see Reviews) should contain an extensive introduction, a 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 also be double-spaced.

Abstract: Authors should provide a brief abstract.

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

1. Jones J. Necrotizing Candida esophagitis.

J A M A .

1980;244:2190-2191.

2. Stryer L.

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e m i s t r y
. 2nd ed. San Francisco, Calif: WH
Freeman Co; 1980:559-596.

Disclosure of Commercial Interests

The 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

GUIDE TO DSM-IV AND ICD-10 CODES

was a big of the Alphaireau Tine With Forth Oracle With Daylor	DSM-IV	ICD-10
ementia of the Alzheimer Type, With Early Onset With Depressed Mood pecify if: With Behavioral Disturbance ementia of the Alzheimer's Type, With Late Onset With Depressed Mood	290.13	F00.03
pecify if: With Behavioral Disturbance	290.21	F00.13
elirium Due to: Indicate General Medical Condition	293.0	F05.0
sychotic Disorder Due to: Indicate General Medical Condition With Delusions	293.81	F06.2
ith Hallucinations	293.82	F06.0
lood Disorder Due to: Indicate General Medical Condition	293.83	F06
nxiety Disorder Due to: Indicate General Medical Condition	293.89	F06.4
mnestic Disorder Due to: Indicate General Medical Condition ementia NOS	294.0 294.8	F02.8 F03
mnestic Disorder NOS	294.8	R41.3
chizophrenia	295	F20
chizophrenia—Disorganized Type	295.10	F20.1
chizophrenia—Catatonic Type	295.20	F20.2
chizophrenia—Paranoid Type	295.30	F20.0
hizophrenia—Residual Type	295.60	F20.5
chizoaffective Disorder	295.70	F25
chizophrenia—Undifferentiated Type	295.90	F20.3
ajor Depressive Disorder	296	F32
polar I Disorder polar Disorder NOS	296 296.80	F30 F39
polar II Disorder	296.89	F31.8
ood Disorder NOS	296.90	F39
sychotic Disorder NOS	298.9	F29
itistic Disorder	299.00	F84
perger's Disorder	299.80	F84.5
rvasive Developmental Disorder NOS	299.80	F84.9
nxiety Disorder NOS	300.00	F41.9
nic Disorder Without Agoraphobia	300.01	F41
eneralized Anxiety Disorder	300.02	F41.1
ssociative Identity Disorder ssociative Disorder NOS	300.14 300.15	F44.81 F44.9
sciolative Disorder NOS	300.19	F68.1
anic Disorder With Agoraphobia	300.19	F40.01
oraphobia Without History of Panic Disorder	300.22	F40.01
ocial Phobia	300.23	F40.1
pecific Phobia	300.29	F40.2
osessive-Compulsive Disorder	300.3	F42.8
ysthymic Disorder	300.4	F34.1
epersonalization Disorder	300.6	F48.1
ody Dysmorphic Disorder	300.7	F45.2
omatization Disorder	300.81	F45.
omatoform Disorder NOS	300.81	F45.9
yclothymic Disorder cohol Dependence	301.13 303.90	F34 F10.2
pocaine Dependence	304.20	F14.2
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cohol Abuse	305.00	F10.1
annabis Abuse	305.20	F12.1
ocaine Abuse	305.60	F14.1
nphetamine Abuse	305.70	F15.1
uttering	307.0	F98.5
orexia Nervosa	307.1	F50
Disorder NOS	307.20	F95.9
urette Disorder	307.23	F95.2
mary Hypercompia	307.42 307.44	F51.0 F51.1
mary Hypersomnia pepwalking Disorder	<u>307.44</u> 307.46	F51.1 F51.3
ssomnia NOS	307.47	F51.3 F51.9
thtmare Disorder	307.47	F51.5
rasomnia NOS	307.47	F51.8
ting Disorder NOS	307.50	F50.9
limia Nervosa	307.51	F50.2
eding Disorders of Infancy or Early Childhood	307.59	F98.2
mmunication Disorder NOS	307.9	F80.9
sttraumatic Stress Disorder	309.81	F43.1
pressive Disorder NOS	311	F32.9
pulse-Control Disorder NOS	312.30	F63.9
thological Gambling	312.31 312.33	F63.0 F63.1
romania eptomania	312.33	F63.1 F63.2
chotillomania	312.39	F63.3
sruptive Behavior Disorder NOS	312.9	F91.9
tention-Deficit/Hyperactivity Disorder, Combined Type	314.01	F90
ention-Deficit/Hyperactivity Disorder NOS	314.9	F90.9
arning Disorder NOS	315.9	F81.9
velopmental Coordination Disorder	315.4	F82
rcolepsy	347	G47.4
	780	G47

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☐ The Black Book of Psychotropic Dosing and Monitoring 2000



Brief Summary

See package insert for full prescribing information.

Indications and Usage. Effector XR is indicated for the treatment of depression and for the treatment of Generalized Annéely Usardo (FGD).

Contraindications: Effector XR is indicated for the treatment of depression and for the treatment of Generalized Annéely Usardo (FGD).

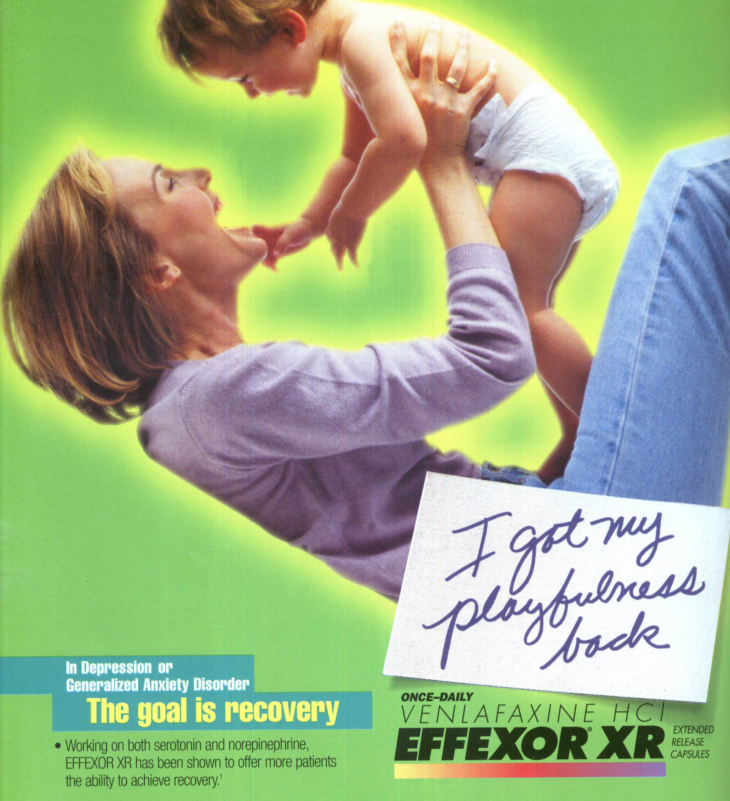
Contraindications: Effector XR is indicated for the treatment to be hypersensitive to veniatazine hydrochloride.

Contraindications: Effector is the contraindicated in patients who have recently been discontinued from an MAOI and started on veniatazine, or who have recently had veniatazine therapy discontinued prior to initiation of an MAOI. These reactions have included tremor, myoclonus, diaphoresis, nausea, womiting, flushing, dizziness, hyperthermia with features resembling neurologic malignant syndrome, seizures, and death. In patients receiving antidepressants with pharmacological properties similar to veniatazine in combination with an MAOI, there have also been reports of serious, sometimes fatal, reactions. For a selective serionin reuptake inhibitor, these reactions have included hyperthermia, flgiffy, myoclonus, autonomic instability with possible regold fluctuations of vital signs, and mental estatuc changes that hotale externed seglication progressing to report thermia and seizures, sometimes fatal, have been reported in association with the combined use of tri-cyclic antidepressants and MAOIs. These reactions have also been reported in patients with bave recently discontinued these drugs and have been started on an MAOI. The effects of combined use of veniatazine and MAOIs have recently discontinued these drugs and have been started on an MAOI. The effects of combined use of veniatazine and MAOIs have recently effective the service of the

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Please see brief summary of Prescribing Information on adjacent page.

The efficacy and safety of EFFEXOR XR for pediatric use have not been established.

EFFEXOR XR is contraindicated in patients taking monoamine oxidase inhibitors (MAOIs). 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 EFFEXOR XR before starting an MAOI.

The most common adverse events reported in EFFEXOR XR placebo-controlled depression trials (incidence ≥10% and ≥2× that of placebo) were nausea, dizziness, somnolence,

Reference: 1. Data on file, Wyeth-Ayerst Laboratories, Philadelphia, Pa.

abnormal ejaculation, sweating, dry mouth, and nervousness; and in GAD trials were nausea, dry mouth, insomnia, abnormal ejaculation, anorexia, constipation, nervousness, and sweating.

Treatment with venlafaxine is associated with sustained increases in blood pressure (BP) in some patients. Three percent of EFFEXOR XR patients in depression studies (doses of 75 to 375 mg/day) and 0.4% in GAD studies (doses of 75 to 225 mg/day) had sustained BP elevations. Less than 1% discontinued treatment because of elevated BP. Regular BP monitoring is recommended.