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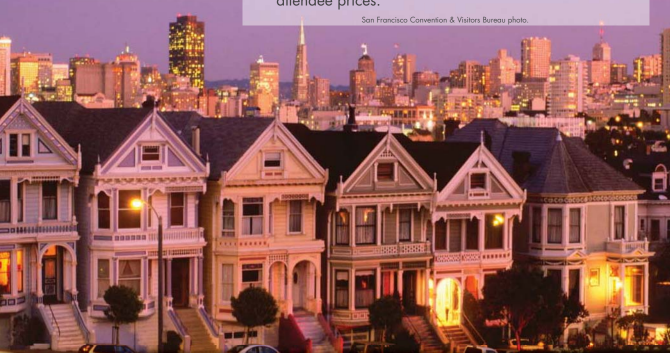
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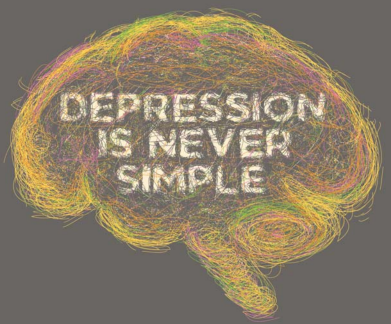
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Venlafaxine Extended Release Tablets (VENLAFAXINE HYDROCHLORIDE)

37.5 mg 75 mg 150 mg 225 mg

IMPORTANT SAFETY INFORMATION

WARNING: Suicidality and Antidepressants
See full Prescribing Information for complete boxed warning.

Increased risk of suicidal thinking and behavior has been reported in children, adolescents and young adults taking antidepressants for major depressive disorder (MDD) and other psychiatric disorders. Venlafaxine Extended Release Tablets are not approved for use in pediatric patients.

Venlafaxine Extended Release Tablets (venlafaxine hydrochloride) are indicated for the treatment of Major Depressive Disorder (MDD) and Social Anxiety Disorder (SAD). Efficacy of venlafaxine HCl was shown in both short-term trials and a longer-term trial in MDD, and in short-term SAD trials. Venlafaxine Extended Release Tablets are contraindicated in patients taking monoamine oxidase inhibitors (MAOIs).

All patients should be monitored appropriately and observed closely for clinical worsening and suicidality, especially at the beginning of drug therapy, or at the time of increases or decreases in dose. Such monitoring should include daily observation by families and caregivers for emergence of agitation, irritability, unusual changes in behavior, or emergence of suicidality.

Venlafaxine Extended Release Tablets 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 Venlafaxine Extended Release Tablets before starting an MAOI. The development of a potentially life-threatening

serotonin syndrome may occur with Venlafaxine Extended Release Tablets, particularly if used concomitantly with serotonergic drugs (including SSRIs, SNRIs, and triptans) or with MAO inhibitors.

Treatment with venlafaxine hydrochloride is associated with sustained hypertension in some patients. Regular blood pressure monitoring is recommended. Mydriasis has been reported in association with venlafaxine; therefore, patients with raised intraocular pressure or those at risk of acute narrow-angle glaucoma should be monitored.

Dosing must be individualized according to the patient's hepatic and renal function status. Abrupt discontinuation or dose reduction has been associated with discontinuation symptoms (generally self-limiting; serious symptoms possible). A gradual reduction in the dose rather than abrupt cessation is recommended.

After treatment with venlafaxine hydrochloride, insomnia and nervousness, activation of mania/hypomania, symptomatic hyponatremia, seizures, abnormal bleeding (most commonly ecchymosis), clinically relevant increases in serum cholesterol, interstitial lung disease and eosinophilic pneumonia have been reported. Venlafaxine Extended Release Tablets should be used cautiously in patients with a history of seizures. Measurement of serum cholesterol should be considered during long-term treatment. Patients should be cautioned about the risk of bleeding associated with concomitant use of Venlafaxine Extended Release Tablets and NSAIDs, aspirin, or other drugs that affect coagulation.

Venlafaxine Extended Release Tablets should be used during pregnancy and nursing only if clearly needed due to the potential for serious adverse reactions.

Adverse reactions occurring in short-term studies of major depressive disorder* were abnormal ejaculation, gastrointestinal complaints (nausea, dry mouth, anorexia), CNS complaints (dizziness, somnolence, abnormal dreams) and sweating. Adverse reactions occurring in short-term studies of social anxiety disorder* were asthenia, gastrointestinal complaints (anorexia, dry mouth, nausea), CNS complaints (anxiety, insomnia, libido decreased, nervousness, somnolence, dizziness), abnormalities of sexual function (abnormal ejaculation, orgasmic dysfunction, impotence), yawn, sweating, and abnormal vision.

*Occurring in at least 5% of patients receiving venlafaxine extended release capsules and at a rate at least twice that of placebo.

Please see brief summary of full Prescribing Information, including complete boxed warning, on adjacent pages.

Reference: 1. Venlafaxine Extended Release Tablets [package insert]. Wilmington, NC: Osmotica Pharmaceutical Corp.; 2008.

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WARNING: SUICIDALITY AND ANTIDEPRESSANT DRUGS

WARNING: SUICIDALITY AND ANTIDEPRESSANT DRUGS
Antidepressants increased the risk compared to placebo of suicidal thinking and behavior
(suicidality) in children, adolescents, and young adults in short-term studies of Major Depressive
Disorder (MDD) and other psychiatric disorders. Anyone considering the use of Venlafaxine
Extended Release Tablets or any other antidepressant in a chilid, adolescent, or young adult must
balance this risk with the clinical need. Short-term studies did not show an increase in the
risk of suicidality with antidepressants compared to placebo in adults beyond age 24; there
was a reduction in risk with antidepressants compared to placebo in adults aged 65 or older.
Depression and certain other psychiatric disorders are themselves associated with increases was a reduction in risk with antidepressants conlipared to piacebo in adults aged of or order.
Depression and certain other psychiatric disorders are themselves associated with increases in the risk of suicide. Patients of all ages who are started on antidepressant therapy should be monitored appropriately and observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. Venlafaxine Extended Release Tablets are not approved for use in pediatric patients. [See Warnings and Precautions and Patient Counseling Information in the full Prescribing Information.] in the full Prescribi ng Information.1

INDICATIONS AND USAGE: Venlafaxine Extended Release Tablets (venlafaxine hydrochloride) are indicated for INDICATIONS AND USAGE: Ventacine Release labels (Ventalist) the hydrocinions) are indicated in the treatment of major depressive disorder (MDD) and Social Anxiety Disorder (SAD), also known as Social Phobia, as defined by DSM-IV. Efficacy of ventafaxine in MDD was shown in both short-term trials and a longer-term trial. Efficacy in SAD was established in short-term trials. CONTRAINDICATIONS: Concomitant use in patients taking monoamine oxidase inhibitors (MAOIs) [see Warnings and Precautions; Potential for interaction with Monoamine Oxidase inhibitors]. WARNINGS AND PRECAUTIONS: Clinical Worsening and Suicide Risk: Patients with MDD, both adult and pediatric, may experience worsening of their depression and/or the emergence of suicidal ideation and behavior (suicidality) or unusual changes in behavior, whether or not they are taking antidepressant. medications, and this risk may persist until significant remission occurs. Suicide is a known risk of depression and other psychiatric disorders, and these disorders themselves are the strongest predictors of suicide. There has been a long-standing concern, however, that antidepressants may have a role in inducing worsening of depression and the emergence of suicidality in certain patients during the early phases of treatment. Pooled analyses of short-term placebo controlled trials of antidepressant drugs (SSRIs and others) showed that these drugs increase the risk of suicidal thinking and behavior (suicidality) in short-term studies in children and adolescents and young adults (ages 18-24) with MDD and other psychiatric disorders. Short-term studies did not show an increase in the adults (alges 16-24) with multiput and other psychiatric disorders. Short-term studies durino show an increase in the risk of suicidality with antidepressants compared to placebo in adults beyond age 24; there was a reduction with antidepressants compared to placebo in adults aged 65 and older. The pooled analyses of placebo-controlled trials in children and adolescents with MDD, obsessive-compulsive disorder, or other psychiatric disorders included a total of 24 short-term trials of 9 antidepressant drugs in over 4400 patients. The pooled analyses of placebo-controlled trials in adults with MDD or other psychiatric disorders included a total of 295 short-term trials (median duration of 2 months) of 11 antidepressant drugs in over 77,000 patients. There was considerable variation in risk of suicidality among drugs but a tendency toward an increase in the volument patients for almost all drugs studied. obtaion of z intomists of 11 antiooptessant origin to et in over 77,000 patients. There was considerable valiation in 11 set of suicidality among drugs, but a tendency toward an increase in the younger patients for almost all drugs studied. There were differences in absolute risk of suicidality across the different indications, with the highest incidence in MDD. The risk differences (drug vs placebo), however, were relatively stable within age strata and across indications. No suicides occurred in any of the pediatric trials. There were suicides in the adult trials, but the number was not sufficient to reach any conclusion about drug effect on suicide. It is unknown whether the suicidality risk extends to longer-term use, i.e., beyond several months. However, there is substantial evidence from placebo-controlled maintenance trials in adults with depression that the use of antidepressants can delay the recurrence of depression. All patients being treated with antidepressants for any indication should be monitored appropriately and observed closely for clinical worsening, suicidality, and unusual changes in behavior, especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases. The following symptoms, anxiety, agitation, panic attacks, insomnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomania, and mania, have been reported in adult and pediatric patients being treated with antidepressants for MDD as well as for other been reported in adout and peduciance patients of several extensions, both psychiatric and nonpsychiatric. Although a causal link between the emergence of such symptoms and either the worsening of depression and/or the emergence of suicidal impulses has not been established, there is concern that such symptoms may represent precursors to emerging suicidality. Consideration should be given to changing the therapeutic regimen, including possibly discontinuing the medication, in patients whose depression is persistently worse, or who are experiencing emergent suicidality or symptoms that might be precursors to worsening depression or suicidality, especially if these symptoms are severe, abrupt in onset, or were not part of the patient's presenting symptoms. If the decision has been made to discontinue treatment, medication should be tapered, as rapidly as is feasible, but with recognition that abrupt discontinuation can be associated with certain symptoms [see Dosage and Administration (2.5) and Warnings and Precautions (5.7) in the full prescribing information for a description of the risks of discontinuation of Venlataxine Extended Felseas Tablets, Families and caregivers of patients being treated with antidepressants for MoD or other indications, both psychiatric and nonpsychiatric, should be alerted about the need to monitor patients for the emergence of agitation, irritability, unusual changes in behavior, and the other symptoms described above, as well as the emergence of suicidality, and to report such symptoms immediately to health care providers. Such monitoring should include daily observation by families and caregivers of Prescriptions for Venlataxine Extended Release Tablets should be written for the smallest quantity of tablets consistent with good patient management, in order to reduce the risk of overdose. Potential for Interaction with monamine Oxidase Inhibitors: Adverse reactions, some serious, have been reported in patients who recently discontinued an MAOI and started on venlataxine hydrochlori indications, both psychiatric and nonpsychiatric. Although a causal link between the emergence of such symptoms With Monoamine Oxidase Inhibitors: Adverse reactions, some serious, have been reported in patients who recently discontinued an MAOI and started on venlafaxine hydrochloride, or who recently discontinued venlafaxine hydrochloride prior to initiation of an MAOI. These reactions included tremor, myoclonus, diaphoresis, nausea, vomiting, flushing, dizziness, hyperthermia with features resembling neuroleptic malignant syndrome, seizures, and death. Venlafaxine Extended Release Tablets 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 venlafaxine hydrochloride before starting an MAOI. A major depressive episode may be the initial presentation of bipolar disorder. It is generally believed (though not established in controlled trials) that treating such an episode with an antidepressant alone may increase the likelihood of precipitation of a mixed/manic episode in patients at risk for bipolar disorder above represent such a conversion is unknown. However, prior to Whether any of the symptoms described above represent such a conversion is unknown. However, prior to initiating treatment with an antidepressant, patients with depressive symptoms should be adequately screened to determine if they are at risk for bipolar disorder. It should be noted that Venlafaxine Evtended Release Tablets are not approved for use in treating bipolar depression. **Serotonin Syndrome**: The development of potentially lifeot approved for use in treating bipolar disports. Serotoniin Syndrome: The development of potentially lifethreatening serotonin syndrome may occur with Venlataxine Extended Release Tablets treatment, particularly with
(1) concomitant use of serotonergic drugs (including SSIRs, SNRIs and triptans) and (2) drugs that impair metabolism of serotonin (see WARNINGS AND
PRECAUTIONS in full Prescribing Information). If concomitant treatment of Venlataxine Extended Release Tablets treatment, particularly with concomitant use of serotonergic drugs (including SSIRs, SNRIs and triptans) and with
drugs that impair metabolism of serotonin (including MAOIs). The concomitant use of Venlataxine Extended
Release Tablets with MAOIs is contraindicated (see Contraindications (4) and Warnings and Precautions (5.2).
Concomitant treatment of Venlataxine Extended Release Tablets with an SSRI, an SNRI, or a 5-hydroxytryptamine
receptor agonist (triptan) is clinically warranted, careful observation of the patient is advised, particularly during
treatment initiation and dose increases. The concomitant use of Venlataxine Extended Release Tablets with
serotonin precursors (such as tryptophan supplements) is not recommended. Sustained Hypertension:
Venlataxine hydrochloride is associated with sustained dose-related increases in blood pressure (BP) in some
patients. Sustained BP increases could have adverse consequences. Postmarketing cases of elevated BP requiring
immediate treatment have been reported. Caution should be exercised in treating patients with pre-existing
hypertension or other underlying conditions that might be compromised by BP increases. Preexisting hypertension or other underlying conditions that might be compromised by BP increases. Preexisting hypertension or other underlying conditions that might be compromised by BP increases. Preexisting PP, perplenicing
sustained increase in BP, either dose reduction or discontinuation should be considered. Elevations in Systolic sustained increase in BP, either dose reduction or discontinuation should be considered. **Elevations in Systolic** and Diastolic Blood Pressure (SBP, DBP): In placebo-controlled premarketing studies, there were changes in mean BP. In most indications, a dose-related increase in SBP and DBP was evident. Across all trials, 1.4% of patients receiving extended-release venlaxafine hydrochloride experienced a ≥15 mm Hg increase in supine DBP with BP \geq 105 mm Hg, compared to 0.9% of patients in the placebo groups. One percent of patients receiving venlaxafine hydrochloride experienced a \geq 20 mm Hg increase in supine SBP with BP \geq 180 mm Hg compared to 0.3% of patients in the placebo groups. **Mydriasis:** Mydriasis has been reported in association with venlafaxine

hydrochloride; patients with raised intraocular pressure or patients at risk for acute narrow-angle glaucoma should be monitored. **Discontinuation of Treatment with Venlafaxine Extended Release Tablets:** Discontinuation symptoms have been systematically evaluated in patients taking venlafaxine, to include prospective analyses of clinical trials and retrospective surveys of trials in MDD and SAD. Abrupt discontinuation or dose reduction of veniafaxina at various doses has been associated with the appearance of new symptoms, the frequency of which increased with increased dose level and longer duration of treatment. Reported symptoms include agitation, anorexia, anxiety, confusion, impaired coordination and balance, diarrhea, dizziness, dry mouth, dysphoric mood, fasciculation, fatique, headaches, hypomania, insomnia, nausea, nervousness, nightmares, sensory disturbances (including shock-like electrical sensations), sommolence, sweating, finitus, tremor, vertigo, and vomiting. During marketing of venlafaxine hydrochloride extended-release capsules, other SNRIs, and SSRIs, there have been spontaneous reports of adverse reactions occurring upon discontinuation of these drugs, particularly when abrupt including the following: dysphoric mood, irritability, agitation, dizziness, sensory disturbances (e.g., paresthesias), anxiety, confusion, headache, lethargy, emotional lability, insomnia, hypomania, tinnitus, and seizures. While these reactions are generally self-limiting, there have been reports of serious discontinuation symptoms. Patiest should be monitored for these symptoms when discontinuing treatment. A gradual reduction in the dose rather than abrupt cessation is recommended whenever possible. If intolerable symptoms occur following a decrease in the dose or upon discontinuation of treatment, then resuming the previously prescribed dose may be considered. Subsequently, the physician may continue decreasing the dose but at a more gradual rate [see Dosage and Administration 2.4 in the Interaction information Incompia and Meryusuness: Treatment-emercinison micrograms. Administration (2.4) in full prescribing information. Insomnia and Nervousness: Treatment-emergent insomnia and nervousness treatment-emergent insomnia and nervousness were more commonly reported for patients treated with venlafaxine hydrochloride extended-release capsules than with placebo in pooled analyses of short-term MDD and other clinical studies, as shown in Table 5 in the full prescribing information. Changes in Weight: In some placebo-controlled trials in MDD, 4% of the patients treated with venlafaxine hydrochloride extended-release capsules and 1% of the placebo-treated patients sustained a loss of 7% or more of body weight during up to 6 months of treatment. The safety and efficacy patients sustained a loss of 7% or more of body weight during up to 6 months of treatment. The safety and efficacy of veniafaxine therapy in combination with weight loss agents have not been established. Co-administration Veniafaxine Extended Release Tablets and weight loss agents is not recommended. Venlafaxine Extended Release Tablets are not indicated for weight loss alone or in combination with other products. **Changes in Height** Rediatric Patients: In the six-month, open-label MDD study, children and adolescents had height increases that were less than expected based on data from age- and sex-matched peers. The difference between observed growth rates and expected growth rates was larger for children (<12 years old) than for adolescents (≥12 years old). **Changes in Appetite**. Adult Patients: Treatment—emergent anorexia was more commonly protect for patients treated with veniafaxine hydrochloride extended-release capsules than for placebo-treated patients in the pool of short-term, double-blind, placebo-controlled MDD (8% vs 4%) and SAD (20% vs 2%) studies. Pediatric Patients: In placebo-controlled trials in MDD and another disorder, 10% of patients aged 6-17 treated with veniafaxine hydrochloride extended-release capsules for up to eight weeks and 3% of patients treated with placebo reported treatment—emergent anorexia. Activation of Mania/Hypomania: Mania rhypomania occurred during MDD studies in 0.3% of patients treated with extended release venlafaxine, the rate was 0.5% compared with 0% of placebo patients. With immediate release venlafaxine, the rate was 0.5% compared with 0% of placebo patients. With immediate release venlafaxine, the rate was 0.5% compared with 0% of placebo patients. With immediate release venlafaxine, the rate was 0.5% compared with 0% of placebo patients. With immediate release venlafaxine, the rate was 0.5% compared with 0% of placebo patients. placebo platents. With infinited lettease verificatine, the face was 0.5% compared with 0% of placebo platents. No reports of mania or hypomania were reported in trials with SAD. As with all drugs effective in the treatment of MDD, Venlafaxine Extended Release Tablets should be used cautiously in patients with a history of mania. Hyponatremia: Hyponatremia may occur as a result of treatment with SSRIs and SNRIs, including Venlafaxine Extended Release Tablets. In many cases, this hyponatremia appears to be the result of the syndrome of inappropriate antidiuretic hormone secretion (SIADH). Cases with serum sodium lower than 110 mmol/L have been reported. Elderly patients may be at greater risk of developing hyponatremia with SSRIs and SNRIs. Also, so states to the control of the control of the Strategy of been reported. Elderly patients may be at greater risk of developing hyponatremia with SSRIs and SNRIs. Also, patients taking diuretics or who are otherwise volumes depleted may be at greater risk [see Use in Specific Populations (8.5) in full prescribing information]. Discontinuation of Ventafaxine Extended Release Tablets should be considered in patients with symptomatic hyponatremia, and appropriate medical intervention should be instituted. Setzures: In all premarketing ventafaxine hydrochloride MDD trials, seizures were reported in 0.3% of ventafaxine hydrochloride-treated patients. Ventafaxine Extended Release Tablets should be used cautiously in patients with a history of seizures and should be discontinued in any patient who develops seizures. Abnormal Bleeding: SSRIs and SNRIs, including Ventafaxine Extended Release Tablets, may increase the risk of bleeding events. Concomitant use of aspirin, nonsteroidal anti-inflammatory drugs (NSAIDs), warfarin, and other anticoagulants may add to this risk. Bleeding events related to SSRIs and SNRIs use have ranged from ecchymoses, hematomas, epistaxis, and petechiae to life-threatening hemorrhages. Patients should be cautioned about the risk of bleeding associated with the concomitant use of Ventafaxine Extended Release Tablets and other drugs that affect coaquidation. Serum Cholesterol Elevation: Clinically relevant increases in serum cholesterol were affect coagulation. Serum Cholesterol Elevation: Clinically relevant increases in serum cholesterol were recorded in 5.3% of ventafaxine hydrochloride-treated patients and 0.0% of patients receiving placebo for at least 3 months in trials. Measurement of serum cholesterol levels should be considered during long-term treatment. Interstitial Lung Disease and Eosinophilic Pneumonia: Interstitial lung disease and eosinophilic pneumonia associated with venlafaxine therapy have been rarely reported. The possibility of these adverse reactions should be considered in venlafaxine-treated patients who present with progressive dyspnea, cough, or chest discomfort. De considered in Venicalaxine-treated patients with present with progressive dyspired, cough; or oriest usscontions. Such patients should undergo prompt medical evaluation, and discontinuation of venidaxine therapy should be considered. **Use in Patients with Heart Disease:** Premarketing experience with venidaxine in patients with concomitant systemic illness is limited. Caution is advised in administering Venlataxine Extended Release Tablets to patients with diseases or conditions that could affect hemodynamic responses. Venidaxine has not been evaluated or used to any appreciable extent in patients with a recent history of myocardial infarction or unstable heart diseases. Patients with these diagnoses were systematically excluded from many clinical studies during venidaxine's premarketing testing. As increases in heart rate (mean increase of 4 beats per minute in MDD trials and 5 heats per minute in SAD trials were observed; caution should be evervised in natients whose underlying underlying the progression of the and 5 beats per minute in SAD trials) were observed, caution should be exercised in patients whose underlying medical conditions might be compromised by increases in heart rate (e.g., patients with hyperthyroidism, heart failure, or recent myocardial infarction). ADVERSE REACTIONS: Clinical Studies Experience: Short-Term, Placebo-Controlled Trials: Adverse Events Leading to Discontinuation of Treatment: Approximately 11% of the 357 patients who received venlafaxine hydrochloride extended-release capsules in MDD trials discontinued treatment due to an adverse reaction (vs 6% of the 285 placebo-treated patients). Adverse reactions that led to treatment discontinuation in at least 2% of drug-treated patients were nausea, dizziness and sommolence. Approximately 17% of the 277 patients in SAD trials who received venlaxafine hydrochloride extended-release capsules discontinued treatment due to an adverse reaction (vs 5% of the 274 placebo-treated patients). Adverse reactions that led to treatment discontinuation in at least 2% of drug-treated patients were nausea, insomnia, reactions that led to treatment discontinuation in at least 2% of drug-treated patients were nausea, insomnia, impotence, headache, dizziness and somnolence. Adverse Events Occurring at an Incidence of 5% or More Major Depressive Disorder: Note in particular the following adverse reactions that occurred in at least 5% of the patients receiving venifataxine hydrochloride extended-release capsules and at a rate at least twice that of the placebo group for all placebo-controlled trials for the MDD indication (see Table 6): Abnormal ejaculation, gastrointestinal complaints (nausea, dry mouth, and anorexia), CNS complaints (dizziness, somnolence, and abnormal dreams), and sweating. In the two U.S. placebo-controlled trials, the following additional reactions occurred in at least 5% of patients treated with venifative hydrochloride extended-release capsules (n = 192) and at a rate at least twice that of the placebo group: Abnormalities of sexual function (impotence in men, anorgasmia in women, and libido decreased), agastrointestinal complaints (constructions) cardiovascular effects (hypertension and vasodilatation), and yawning. Social Anxiety Disorder: Note in particular the following adverse reactions that occurred in at least 5% of the patients receiving evalidaxine hydrochloride extended-release capsules and at a rate at least twice that of the placebo group for the 2 placebo-controlled trials for the SAD indication (see Table 7): Asthenia, qastrointestinal complaints (anorexia, constitution, dry mouth, nausea), release capsules and at a rate at least wice that of the placebor group to the 2 placebor-Common tank of the RAD indication (see Table 7). Asthenia, gastrointestinal complaints (anorexia, constipation, dry mouth, nausea), CNS complaints (dizziness, insomnia, libido decreased, nervousness, somnolence), abnormalities of sexual function (abnormal ejaculation, impotence, libido decreased, orgasmic dystunction), yawn, sweating, and abnormal vision. Adverse Events Occurring at an Incidence of 2% or More: MDD and SAD trials included patients receiving venlafaxine hydrochloride extended-release capsules in doses ranging from 75 mg to 225 mg/day for up to 12 weeks. The prescriber should be aware that the following adverse reactions figures cannot be used to receive the venture of the prescriber of the curves of the question of the prescriber of the curves of the question of the prescriber of the properties. predict the incidence of adverse reactions in the course of usual medical practice. Similarly, the cited frequencies predict the includer or adverse reactions in the course or usual medical practice. Similarly, include the capacitors cannot be compared with figures obtained from other clinical investigations involving different treatments, uses, and investigators. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and nondrug factors to adverse reaction incidence rate in the population studied. [See TABLE 6 in full Prescribing Information.] TABLE 6: Treatment Temergent Adverse Reaction Incidence in Short-Term Placebo-Controlled Clinical Trials with Venlafaxine Hydrochloride Extended-Release Capsules in Patients with Major Depressive Disorder. This table reports adverse events that occurred in 2% of the property of the proper or more of patients treated with venidaxine hydrochioride extended-release capsules where the incidence in patients treated with venidaxine hydrochioride extended-release capsules (n=357) was greater than the incidence for the respective placebo-treated patients (n=285). For each adverse reaction, the incidence of reactions in the drug-treated patients is listed before the incidence in placebo-treated patients. **Body as a Whole:** Asthenia (8% and 7%). **Cardiovascular System:** Vascolitation (4% and 2%); Hypertension (4% and 1%). **Digestive System:** Aussea (31% and 7%); Constituation (8% and 5%); Anorexia (8% and 4%); Vomitting (4% and 2%); Hallence (4% and 3%). **Metabolic/Nutritional:** Weight Loss (3% and 0%). **Nervous System:** Dizziness (20% and 9%);

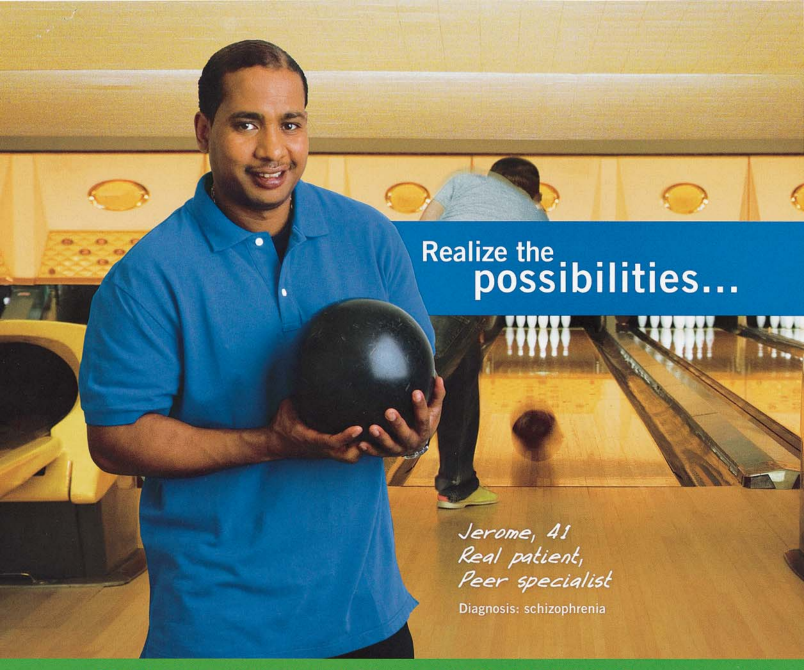
Somnolence (17% and 8%) Insomnia (17% and 11%); Dry mouth (12% and 6%); Nervousness (10% and 5%); Somnolence (17% and 8%) Insomnia (17% and 11%); Dry mouth (12% and 6%); Nervousness (10% and 5%); Abnormal Dreams (7% and 2%); Tremor (5% and 2%); Depression (3% and <1%); Persethseia (3% and 1%); Libido Decreased (3% and <1%); Agitation (3% and 1%). Respiratory System: Pharyngitis (7% and 6%); Yawn (3% and 0%). Skin: Sweating (14% and 3%). Special Senses: Abnormal vision (4% and <1%). Urogenital System: Abnormal ejaculation (16% and <1%); Impotence (4% and <1%); Female anorgasmia (3% and <1%). [See TABLE 7 in full Prescribing Information]. TABLE 7: Treatment Emergent Adverse Reaction Incidence in Short-Term Placebo-Controlled Clinical Trials with Venlafaxine Hydrochloride Extended-Release Capsules in Patients with Social Anxiety Disorder. This table reports adverse events that occurred in 2% or more of patients treated with venlafaxine hydrochloride extended-release capsules where the incidence for the respective placebo-treated patients (n=274). For each adverse reaction, the incidence of reactions in the drug-treated natients is listed hefore; the incidence in placebo-treated patients (n=274). For each adverse reaction, the incidence of reactions in the drug-treated natients is listed hefore; the incidence in placebo-treated patients. Body as a Whole: Neadache (34%). treated with venlafaxine hydrochloride extended-release capsules (n=277) was greater than the incidence for the espective placebo-treated patients in E-274). For each adverse reaction, the incidence of reactions in the drugtered patients is listed before the incidence in placebo-treated patients. Body as a Whole: Headache (34% and 33%): Asthenia (17% and 8%); Flu Syndrome (6% and 5%); Accidental Injury (5% and 3%); Abdominal Paid (4% and 3%). Cardiovascular System: Hypertension (6% and 5%); Ascoliation (3% and 1%); Palpitation (3% and 1%). Digestive System: Nausea (29% and 9%); Anorexia (20% and 1%); Constipation (8% and 4%); Diarrhea (6% and 5%); Vomiting (3% and 2%); Eructation (2% and 0%). Metabulic/Mutritional: Weight Loss (4% and 5%); Parest (5% and 3%); Elbido Decreased (9% and -1%); Dizziness (16% and 8%); Somnolence (16% and 8%); Nervousness (11% and 3%); Libido Decreased (9% and -1%); Anxiety (5% and 3%); Twitching (2% and 0%). Respiratory System: Yawn (5% and <1%); Sinusitis (2% and 1%); Skin: Sweating (13% and 2%). Special Senses: Abnormal vision (6% and 3%). Urogental System: Abnormal ejaculation (16% and 1%); Impotence (10% and 1%); Female Orgasmic Dysfunction (8% and 0%). Vital Sign Changes: Venlafaxine hydrochloride was associated with a mean increase in pulse rate of 4 beats/min in SAD trials. In premarketing trials, the mean change from baseline heart rate for patients treated with extended-release venlafaxine hydrochloride in MDD and SAD trials was 4 beats-per-minute and 5 beats-per-minute, respectively. In a flexible-ose study with doses ranging from 200 mg to 375 mg/day, patients receiving extended-release venlafaxine hydrochloride had a mean increase in heart rate of 8.5 beats-per-minute, respectively. In a flexible-ose study with doses of venlafaxine hydrochloride immediate-release tablets in the rang EXPERIENCE: Voluntary reports or other adverse reactions temporarily associated with rise use of vertilatable her received since market introduction. Because these reactions have been reported from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure. These reports include the following reactions: agranulocytosis, anaphylaxis, aplastic anemia, catatonia, congenital anomalies, impaired coordination and balance, CPK increased, deep vein thrombophlebitis, delirium, EKG abnormalities such as CT protongation; cardiac arrhythmias including atrial fibrillation, supraventricular tachycardia, ventricular extrasystoles, and rare reports of ventricular fibrillation and ventroular tachycardia, including torsade de pointes; epidermal necrolysis/Stevens-Johnson syndrome, erythema multiforme, extraveryandial emmons; fincluding foundations discussions and tardies deviated since deviated and control extrapyramidal symptoms (including dyskinesia and tardive dyskinesia), angle-closure glaucoma, hemorrhage (including eye and gastrointestinal bleeding), hepatic reactions (including GGT elevation; abnormalities of unspecified liver function tests; liver damage, necrosis, or failure; and fatty liver), interstitial lung disease, involuntary movements, LDH increased, neuroleptic malignant syndrome-like reactions (including a case of a 10-year-old who may have been taking methylphenidate, was treated and recovered), neutropenia, night sweats, pancreatitis, pancytopenia, panic, prolactin increased, renal failure, rhabdomyolysis, serotonin syndrome, shocklike electrical sensations or tinnitus (in some cases, subsequent to the discontinuation of venlataxine or tapering of dose), and syndrome of inappropriate antidiuretic hormone secretion (usually in the elderly). **DRUG** INTERACTIONS: Alcohol: The effect of alcohol on plasma levels of Venlataxine Extended Release Tablets is not INTERACTIONS: Alcohol: The effect of alcohol on plasma levels of Ventafaxine Extended Release Tablets is not known. Cimetidine: Use caution when administering ventafaxine hydrochloride with cimetidine to patients with preexisting hypertension or hepatic dysfunction, and the elderly. Diazepam: A single dose of diazepam did not appear to affect the PK of either ventafaxine hydrochloride (150 mg/day) or its major active metabolite, O-desmethylventafaxine (DVD). Ventafaxine hydrochloride (150 mg/day) or the PK of diazepam or its active metabolite, desmethyldiazepam, or affect the psychomotor and psychometric effects induced by diazepam. Haloperidol: Ventafaxine hydrochloride (150 mg/day) decreased total oral-dose clearance of haloperidol resulting in a Ventafaxine hydrochloride (150 mg/day) decreased total ovate the haloperidol elimination trus use unchanged. Lithium: A single dose of lithium (600 mg) did not appear to affect the PK of either ventafaxine hydrochloride (150 mg/day) der optional proteins: ventafaxine hydrochloride is not highly bound to plasma proteins; coadministration of Ventafaxine Extended Release Tablets and a highly protein-bound drug should not cause increased free concentrations of the other drug. Drugs That Inhibit (Votochrome P450 Isenzymes: CYP206 and CYP3A4 Inhibitors: Ventafaxine hydrochloride is metabolized to ODV by CYP206. Drugs inhibiting this isoenzyme have the potential to increase plasma concentrations of ventafaxine hydrochloride and ODV are approximately equiactive and equipotent, no dosage adjustment is required venlafaxine hydrochloride and ODV are approximately equiactive and equipotent, no dosage adjustment is required when venlafaxine hydrochloride is coadministered with a CYP2D6 inhibitor. Pharmacokinetic studies with ketoconazole in both poor and extensive metabolizers of CYP2D6 resulted in higher plasma concentrations and when venlafaxine hydrochloride is coadministered with a CYP2D6 inhibitor. Pharmacokinetic studies with ketoconazole in both poor and extensive metabolizers of CYP2D6 resulted in higher plasma concentrations and AUCs of both venlafaxine hydrochloride and ODV in most subjects following administration of ketoconazole. Concomitant use of CYP2D6 resulted in higher plasma concentrations and AUCs of both venlafaxine hydrochloride and ODV. Use caution if therapy includes venlafaxine hydrochloride and any CYP3A4 inhibitors and venlafaxine hydrochloride and any CYP3A4 inhibitor. Drugs Metabolized by Cytochrome P450 isoenzymes: Venlafaxine hydrochloride is a relatively weak inhibitor of CYP2D6 in vitro. Impramine: Venlafaxine hydrochloride did not affect the PK6 filimpramine or CYP3A4 inhibitor. The constraint in vitro. Impramine: Venlafaxine hydrochloride dose of venlafaxine hydrochlorid for serotoni syndrome, caution is advised when Venlafaxine Extended Release Tablets are coadministered with other drugs that may affect the serotonergic neurotransmitter systems, such as triptans, SSRIs, other SNRIs, linezolid, lithium, tramadol, or St. John's Wort. If concomitant treatment of Venlafaxine Extended Release Tablets with these drugs is warranted, careful observation of the patient is advised, particularly during treatment initiation and dose increases. Concomitant use of Venlafaxine Extended Release Tablets with tryptophan supplements is not recommended [see WARNINGS AND PRECAUTIONS in full Prescribing Information]. There have been rare recommended [see WARNINGS AND PRECAUTIONS in full Prescribing Information]. There have been rare postmarketing reports of serotonin syndrome with use of an SSRI and a triptan. If concomitant use of Venlafaxine Hydrochloride Extended Release tablets with a triptan is warranted, careful observation of the patient is advised, particularly during treatment initiation and dose increases [see WARNINGS AND PRECAUTIONS in (Hzerscribing) Information]. Drugs That Interfere With Hemostasis: Interference with serotonin reuptake may affect platelet function and result in bleeding. Concurrent use of NSAIDs or aspirin may increase this risk. Increases in prothrombin time (PT), partial thromboplastin time (PT), or INR have been reported when venladavine Hydrochloride was given to patients on warfarin should be carefully monitored when Venlafaxine Extended Release Tablets are begun or discontinued. Electroconvulsive Therapy: There is no clinical data establishing the benefit of electroconvulsive therapy combined with Venlafaxine Hydrochloride Extended Release Tablets. Postmarketing Spontaneous Drug Interaction Reports: There have been reports of elevated clozapine levels temporally associated with adverse reactions, including seizures, following the addition of venlaxafine. There have been reports of increases in PT.PT, or INR when venlafaxine as given to patients also receiving warfarin. USE IN SPECIFIC POPULATIONS: Pregnancy: Teratogenic Effects: Pregnancy Category C: There are no adequate and well-controlled studies of venlafaxine in pregnant women.

Venlafaxine Extended Release Tablets should be used during pregnancy only if clearly needed. Non-Teratogenic Effects: Neonates exposed to venlafaxine hydrochloride late in the third trimester have developed compilications requiring prolonged hospitalization, respiratory distress, syanosis, apnea, seizures, unstable temperature, feeding difficulty, worlting, hypogycemia, hypo- and hypertonia, hyperrellexia, tremor, litteriness, irritability, and constant crying. This is consistent with a toxic effect of SSRIs or SNRIs or a drug discontinuation syndrome. In some cases, it is consistent with a toxic effect of SSRIs or SNRIs or a drug discontinuation syndrome. In some cases, it is consistent with servotion syndrome. When treating a pregnant woman with Venlafaxine Extended Release Tablets during the third trimester, carefully consider the potential risks and benefits of treatment. Labor and Delivery: The effect of venlafaxine hydrochloride on labor and delivery in humans is unknown. Nursing Mothers: Venlafaxine hydrochloride and ODV, its active metabolite, are excreted in human milk. Because of the potential for serious adverse reactions in nursing infants, a decision should be made whether to discontinue of the decision in unsing infants, a decision should be made whether to discontinue ventafaxine Extended Release Tablets, taking into account the importance of the drug to the mother. Pediatric Use: Safety and effectiveness in the pediatric population have not been established [see BOXED WARNING and Warnings and Precautions: Clinical Worsening and Suicide Risk]. Anyone considering using Venlafaxine Extended Release Tablets in a child or adolescent must balance the potential risks with the clinical need. While no studies have adequately assessed the impact of venlafaxine hydrochloride on growth, development, and maturation of children and adolescents, studies suggest it may adversely affect weight and height is recommended with total may adversely affect weight and height is recommended during treatment, particularl

To report SUSPECTED ADVERSE REACTIONS, contact Upstate Pharma, LLC Pharmaceutical Corp. at 1-888-299-1053 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

This brief summary is based on Venlafaxine Extended Release Tablets Prescribing Information, August 2008. Osmotica Pharmaceutical Corp.

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- Effectively treats the symptoms of schizophrenia
- Well-established tolerability profile

GEODON is indicated for the treatment of schizophrenia.

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.

Individual results may vary.

Please see brief summary of prescribing information on adjacent page.

For more information, please visit www.pfizerpro.com/GEODON

■ Target 120–160 mg/day with meals

-initiate at 40 mg/day

—lowest effective dose should be used

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 short-term schizophrenia trials, the most commonly observed adverse events associated with GEODON at an incidence of ≥5% and at least twice the rate of placebo were somnolence and respiratory tract infection.

In short-term schizophrenia clinical trials, 10% of GEODON-treated patients experienced a weight gain of ≥7% of body weight vs 4% for placebo.



Increased Mortality in Elderly Patients with Dementia-Related Psychosis: Elderly patients with dementia-related psychosis breated 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 of between 1.6 to 1.7 times that seen in placebo-treated patients. Over the course of a typical 10 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 deaths appeared to be either cardiovascular (e.g., heart failure, sudden each) 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 schizophrenia and acute manic or mixed episodes associated with pipolar disorder.

Schroupitents passers — *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 myocardial infarction, or with uncompensated hear failure (see WARNINGS). Pharmacokinetic/pharmacodynamic studies between GEODON and other drugs that prolong the QT interval have not been with MININGS.) Primace Sonder Districts of CEDON and the drough that prolong the OT interval cannot be enclosed. Therefore, GEDON the should not be given with ordinative, solds of membraconsolvanies studies between GEDON and other during that prolong the OT interval cannot be enclosed. Therefore, GEDON is should not be given with ordinative, solds of membraconsolvanies, membraconsolvanies, membraconsolvanies, and the providence of the provi performed. An additive effect of GEODON and other drugs that protong the OT interval cannot be excluded. Therefore, ECOON should not be given with detellide, sotalol, quinidine, other Class la and Ill artistrythmics, mesoridazine, thioridazine, chlorpromazine, droperidol, pimozide, sparfloxacin, gatifloxacin, moxilloxacin, halofantrine, mefloquine, pentamidine, arsenic trioxide, levomethadyl acetate, dolasetron 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, hypoteolenia, treatment with antihypertensive medications). Segürzes; in clinical trials, seziurze socurred in 0.4% of GEODOM patients. There were confounding factors that may have contributed to seizures in many of these cases. As with other antipsychotic drugs, GEODON should be condumning factors that may have continuous to secures in many or these cases. As winn or employconic units, economic used cautiously in patients with a history of secures or with conditions that potentially lower the secure threshold. e.g., Atheriner's dementia. Conditions that lower the seizure threshold may be more prevalent in a population of 65 years or older. <u>Dysphagia</u>, Esophageal dysmotility and aspiration have been associated with antipsychold circula use. Aspiration pneumonia is a common cause of morbidity and morbality and delicity patients, in particular those with advanced Atheriner's dementia, and ECOOM and other antipsychold circular should be used cautiously in patients at risk for aspiration pneumonia. (See also Boxed WARNING, WARNINGS: Increased Mortality in Elderty Patients with Dementia-Related Psychosis, hyperprolactinemia; As with other drugs that antagonize dopamine D, receptors, ECDOON elevates prolactin levels in humans. Tissue culture experiments indicate that approximately one third of human breast 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 tumorigeness in humans; the available evidence is considered too limited to be conclusive at this time. <u>Potential for Cognition and Motor Impairment</u>, Somnolence was a commonly reported adverse event in GEODOM patients. In the 4- and 6-week placebo-controlled trials, somnolence was reported in 14% of GEODOM patients in the 4- and 6-week placebo-controlled trials, somnolence was reported in 14% of GEODOM patients in the 3- and 4- and 5- and trials, somnolence was reported in 14% of GEODON patients vs 7% of placebo patients. Somnolence led to discontinuation in 0.3% of patients in short-term clinical trials. Since GEODON patients vs 7% of placebo patients. Somnolence led to discontinuation in 0.3% of patients in short-term clinical trials. Since GEODON has the potential of in inpair judgment, thinking, or motor skills, patients should be cautioned about performing activities requiring mental alertness, such as operating a motor vehicle (including automobiles) or operating hazardous machinery until they are reasonably certain that GEODON therapy does not affect them adversely. Pringism: One case of pringism was reported in the premarketing database. Body Temperature Regulation: Although not reported with GEODON immensiteling trials, disruption of the body's ability to reduce core body temperature Regulation: Although not reported with GEODON immensiteling trials, disruption of the body's ability to reduce core body temperature has been attributed to antipsychotic agents. Suicide: The possibility of a suicide attempt is inherent in psychotic illness and close supervision of high-risk patients should accompany drug therapy. GEODON prescriptions should be written for the smallest quantity of capsules consistent with good patient management to remarketing clinical studies. Because of the risk of OT, prolongation and orthostatic hypotension with GEODON (aution should be observed in cardiac patients) see all Prolongation and Risk of Sudden Death in WARNINGS and Othostatic Hypotension in PRECUTIONS). 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

surements. Low serum potassium and magnesium should be repleted before treatment. Patients who are started on diuretics during measurements. Low serum potassium and magnesium should be repleted before treatment. Patients who are started on diureties during GEODON therapy need periodic monitoring of serum potassium and magnesium. Discontinue GEODON in patients who are found to have persistent QT, measurements >500 msec (see WARNINGS). Drug Interactions: (1) GEODON should not be used with any drug that prolongs the QT interval. (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 levodopa and dopamine agonists. Effect of Other Drugs on GEODON: Cartamazepine, 200 mg bid for 21 days, resulted in a decrease of approximately 35% in the AUC or GEODON. Retocorazule, a potent inhibitor of CYP3A4, 400 mg of for 5 days, increased the AUC and C_{my} of GEODON by about 35% -40%. Certification, 800 mg qif for 2 days, did not affect GEODON pharmacokinetics. Coadministration of 30 mL of Maakovidi not affect GEODON pharmacokinetics. Population pharmacokinetic analysis of achievablement positions in the properties of the properties of the properties of the part analysis of achievablement centre in the properties of of schizophrenic patients in controlled clinical trials has not revealed any clinically significant pharmacoline interactions of schizophrenic patients in controlled clinical trials has not revealed any clinically significant pharmacoline interactions propranolol, or forazepam. Effect of SCDON on Other Drugs; in vitro studies revealed little potential for GEODON to interfere with the metabolism of drugs cleared primarily by CYP1A2, CYP2O9, CYP2O19, CYP2O19, and CYP3A4, and little potential for drug interactions with GEODON to ted significant in GEODON 40 mg bid administered concomitantly with thinium 450 mg bid for 7 days did not affect the steady-Security due to displacement, security 4 mit good annimisered concominatiny with inhumination in ground in due to a state level or renal clearance of lithium. GEODON 20 mp bid din or affect the pharmacokinetics of concominatiny administered oral contraceptives, ethiny fest addition, and even orgester (0.15 mg). Consistent with in vitro results, a study in normal healthy voluntees showed that GEODON did not alter the relabolism of deutromethynian, a CYP206 model substrate, to its major metabolist, edutrophan. There was no statistically significant change in the urinary dextromethorphan/dextrorphan ratio. Carcinogenesis, Mutagenesis, Impairment of Fertility: Lifetime carcinogenicity studies were conducted with GEODON in Long Evans ratis and DO-1 mice. In male mice there was no increase in incidence of tumors relative to controls. In female mice there were dose-related increases in the incidences of pituliary gland adenorm and carcinoma, and mammany gland adenorma desertion and all doses lested. Increases in serum probactive more cheened in a 1-month distant softwin in female hip that productivin in service in a 5-week diletary. observed in a 1-month dietary study in female, but not male, mice, GEODON had no effect on serum prolactin in rats in a 5-week dietary in one strain of *S. typhimurium* in the absence of metabolic activation. Positive results were obtained in both the in vitro mammalian call gine mutation assay and the in vitro chromosomal abertain assay in human lymphocytes. Impairment of Fertility, GEODON increased gine to copulation in Sprague-Daviley ratis in two fertility and early embryonic development studies at doese of 10 to 160 mg/kg/day (0.5 to 8 times the MRHO of 200 mg/day on a mg/m² basis). Fertility rate was reduced at 160 mg/kg/day (8 times the MRHO on a mg/m² basis). There was no effect on fertility at 40 mg/kg/day (2 times the MRHO on a mg/m² basis). The fertility of fermale rats was reduced. *Pregnancy—Pregnancy—Category C*: There are no adequate and well-controlled studies in pregnant women. SECDON should be used during pregnancy—only if the potential benefit justifies the potential risk to the fetus. *Labor and Delivery*: The effect of GEODON on the delivery in human six unknown. *Nursing Mediters*: this not known whether, and if so in what amount, GEODON or its metabolities are excreted in human milk. It is recommended that women receiving GEODON should not breast feed. *Padiatric Use*: The safety and effectiveness of GEODON in pediatric patients have not been established. *Gertafric Use*: Of the approximately 4500 patients breated with GEODON in clinical studies and the second of the properties of the control of any different tolerability for GEODON or feedood clearance of GEODON in the elderly compared to younger adults. Nevertheless, the presence of multiple factors that might increase the pharmacodynamic response to GEODON or cause poorer tolerance or orthostasis, should lead to consideration of a lower starfing dose, sowerthation, and careful monotring during the initial dosing period for some elderly patients. ADVERSE FRACTIONS—"Adverse Findings pharmacodynamic response to GEODON, or cause poorer loterance or orthoctasis, should lead to consideration of a lower starting observed in an and careful monitoring during the initial dosing period for some elderly patients. AUVERSE REACTIONS — Adverse Findings Observed in Short-term, Placebo-Controlled Trials: The following findings are based on the short-term placebo-controlled premarketing trials for schizophrenia (a pool of two 6-week and two 4-week fixed-dose trials) and bipolar mania (a pool of two 3-week flexible-dose trials) in which ECDOON was administered in doses ranging from 10 to 200 mg/dga, Averse Events Associated with Discontinuation: Schizophrenia: Approximately 4.1% (29/702) of GEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with about 2.2% (6/273) on placebo. The most common event associated with dropout the train including 2 dropouts for rash among GEODON treated patients in short-term, placebo-controlled studies discontinued treatment and approximately 5.5% (18/279) of GEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to an adverse event, compared with short place to a placebo-controlled studies discontinued treatment due to an adverse event compared with short placed patients in short-term, placebo-controlled studies discontinued treatment due to a devenament of the CEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to a devenament of the CEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to a devenament of the CEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to a devenament devenament and the proportion the CEODON-treated patients in short-term, placebo-controlled studies discontinued treatment due to a devenament and the controlled studies discontinued treatment due to a devenament and the controlled studies discontinued treatment due to a devena adverse event, compared with about 3.7% (5/136) on placebo. The most common events associated with dropout in the GEODON-treated patients were alkathisia, anxiety, depression, dizziness, dystonia, rash and vomiting, with 2 dropouts for each of these events among GEODON patients (1%) compared to one placebo patient each for dystonia and rash (1%) and no placebo patients for the remaining adverse events. Adverse Events at an incidence 2-5% and at Least Twice the Rate of Placebo. The most commonly observed adverse events associated Adverse Events at an Incidence ≥5% and at Least Twice the Rate of Placebo: The most commonly observed adverse events associated with GEODON in schiop/brenial trials were somnolence (14%) and respiratory tract infection (%%). The most commonly observed adverse events associated with the use of GEODON in policy of GEODON in policy of a main trials were somnolence (14%), extrapyramidal symptoms (34%), diziness (16%), alactihisia (10%), abnormal vision (6%), ashlenia (6%), and vomiting (5%). The following list enumerates the treatment-emergent of the properties of the control of the cont mouth, increased salivation, arthralgia, arwiely, dizziness, dystonia, hypertonia, somnolence, tremor, rihinitis, rash, and abnormal vision.

Extrapyramidal Symptoms (EPS): The incidence of reported EPS for ECDON patients in the short-term, placebo-controlled schizophrenia irritals was 14% set % for placebo-Objectively collected data from those trails was 14% simpson-Angus Raing Scale and the sess-Authisia Scale did not generally show a difference between GEODON and placebo. Dystonia: Prolonged abnormal contractions of muscle groups may occur in susceptible individuals during first few days of treatment. Dystonia may occur at any dose level but with greater frequency and severity with high potency and at higher doses of first generation antipsychotic drugs. Elevated risk is observed in males and younger age groups. Vital Sign Changes: GEODON is associated with orthostatic hypotension (see PRECAUTIONS). Weight Gain: In short-term groups. Vital sign Changes: GEODON is associated with orthostatic hypotension (see PRECAUTIONS). Weight Gain: in short-term schizophrenia trials, the proportions of patients meeting a weight gain criterion of 2.7% of body weight were compared, revealing satistically significantly greater incidence of weight gain for GEODON patients (10%) vs placebo patients (4%). A median weight gain of 0.5 kg was observed in GEODON patients vs 0.0 kg in placebo patients. Weight gain was reported as an adverse event in 0.4% of both GEODON and placebo patients. During long-term therapy with GEODON, a categorization of patients at baseline on the basis of body mass index (BMI) showed the greatest mean weight gain and the highest incidence of clinically significant weight gain (27% of body weight) in patients with a low BMI (23) compared to normal (23-27) or overweight (27) patients. There was a mean weight gain of 1.4 kg for patients with a "bormal" BMI, and a 1.3 kg mean weight loss for patients with a "high" BMI. CEG Changes:
GEODON is associated with an increase in the OT, interval (see WARNINGS). In schizophrenia trials, GEODON was associated with a mean increase in heart rate of 1.4 beats per minute compared to a 0.2 beats per minute decrease among placebo patients. Other Adverse Eventual of Mean and the Patients of 1.4 Minute of the patients of the increase in heart rate of 1.4 beats per minute compared to a 0.2 beats per minute decrease among placebo patients. Other Adverse Events Observed During the Premarketing Evaluation of GEODON: Frequent adverse events are those occurring in at least 1/100 patients; infrequent adverse events are those occurring in lever than 1/1000 patients. Schizophrenia: Body as a Whotle — Frequent: abdominal pain, flu syndrome, fever, accidental fall, face edema, chills, photoesnistivity reaction, flank pain, hypothermia, motor vehicle accident. <u>Eardiovascular System — Frequent Landyavardia</u>, hyperton, postural hypotension, furfrequent bradycardia, angina pectoris, atrial fibrillation; Rare: first-degree AV block, bundle branch block, philebitis, pulmonary embolus, cardiomeagly, crebral infarct, cerebrovascular accident, deep thrombophilebitis, myocarditis, furrombophiebitis. <u>Digestive</u> system — Frequent Landy, crebral mining, furfrequent rectal themorrhage, dysphagia, tongue edema; Rare gunh emorrhage in minación, gamma glutamy transpeptidase increased, hematemesis, cholestatic jaundice, legal impaction, gamma glutamy transpeptidase increased, hematemesis, cholestatic jaundice, legal in un posticio. Page a biocardo. Page a biocardo de la morria de Impaction, gaining glutain y tallspeptiouse increasor, increasor, increasor, and increasor, and increasor, and a staff were deposit, melena, Engloring—Rare hypothyroidism, Hy thirst, transaminase increased, peripheral edema, hyperglycemia, reatine phosphokinase increased, alkaline phosphatase increased, hypercholesteremia, dehydration, lactic dehydrogenase increased, albuminuria, hypokalemia, Raze BUN increased, creatinine increased, hypertilemia, hypocholesteremia, hy Musculoskeletal System — Frequent: myaliga: Infrequent:tenosynovitis; Rare: myopathy. Nervous System — Frequent: agitation, extrapyramidal syndrome, tremor, dystonia, hypertonia, dyskinesia, hostility, twitching, paresthesia, confusion, vertigo, hypokinesia, admoral gait, oculogyric crisis; hypertenisa, adva, amnesia, cogyweler ligidity, delimin, hypotonia, adviseia, dysarthina, withdrawal syndrome, buccoglossal syndrome, choreoathetosis, diplopia, incoordination, neuropathy; Infrequent: paralysis; Rare: myoclonus, nystagmus, torticollis, circumoral paresthesia, opisthotonos, reflexes increased, trismus. Respiratory System haze. In productivis, in seguinos, torticomo a para seriesa, opisacionos, se inexes intereses o interesente interesente preumonia, epistas, Rare hemophysis, laryngismus, Saha and Appendages — Infraquent maculopapular rash, urticaria, alopecia, eczema, exfoliative dermatitis, contact dermatitis, vesiculobullous rash. Special Senses — Frequent fungal dermatitis, infrequent conjunctivitis, dry eyes, tinnitus, blepharitis, cataract, photophobia; Rare: eye hemorrhage, visual field defect, keratitis, keratoconjunctivitis. <u>Urogenital System</u>—Infraquent: impotence, abnormal ejaculation, amenorrhea, hematuria, menorrhagia, female lactation, polyuria, urinary retention, metrorrhagia, male sexual dystlunction, anongasmia, glycosuria; Rare; gynecomastia, vagite hemorrhage, noturia, oligiuria, female sexual dystlunction, uterine hemorrhage, adverse Indiaglo Boservel in Trials of Inframuscular GEODON: in these studies, the most commonly observed adverse events associated with the use of intramuscular GEODON (25%) and observed at a rate on intramuscular GEODON (in the higher dose groups) at least twice that of the lowest intramuscular GEODON group were observed at a rate on intramuscular GEOODN (in the higher dose groups) at least twice that of the lowest intramuscular GEOODN group were headache (13%), nausea (12%), and somnolence (20%). Adverse Events at an Incidence -1% in Short-Term Fixed-Dose Intramuscular Trials: The following list enumerates the treatment-emergent adverse events that occurred in ≥1% of GEOON patients (in the higher dose groups) and at least twice that of the lowest intramuscular GEOON group. Body as a Whole—headache, injection site pain, astemba addominal pain, its syndrome, back pain. Cardiovascular—postural hypotension, hypertension, bradycardia, vasodilation. Digestine—nausea, rectal hemorrhage, diarrhea, vomiting, dyspepsia, anorexia, constipation, tooth disorder, dry mouth. Nervous—dizzness, anxiety, insomnia, somnolence, akathisia, apitation, extrapyramidal syndrome, hypertonia, cogwhele rigidity, paresthesia, personality disorder, psychosis, speech disorder. Repsidatory—minitis. Skianard Appendages—frurunculosis, sweating. Urgogental—dysemorrhea, priapism.

DRUG ABUSE AND DEPENDENCE—Controlled Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Overland Substance Class: GEODON is not a controlled substance. Ove DROG ABUSE AND DEPENDENCE—Commonles Substance Crass; GEODOW is not a controller substance. DVENDOSABE—In premarketing trials in over 5400 patients, accidental or intentional overdosage of GEODON was documented in 10 patients. All patients survived without sequelae. In the patient taking the largest confirmed amount (3240 mg), the only symptoms reported were minimal sedation, slurring of speech, and transitory hypertension (BP 20095).



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Beth Israel Deaconess Medical Center Harvard Medical School

Beth Israel Deaconess Medical Center (BIDMC) and Harvard Medical School (HMS) are seeking nominations for and applications from an established academic leader with demonstrated clinical, research, and teaching excellence, who will be responsible for leading a distinguished academic Department of Psychiatry. The Beth Israel Deaconess Medical Center, a 665-bed tertiary and quaternary care hospital, is a founding member of CareGroupSM, an organized system of quality healthcare serving individuals, families and communities in New England. The incumbent will be the Bullard Professor of Psychiatry at Harvard Medical School, and will sit on the Executive Committee and Board of Directors of Harvard Medical Faculty Physicians Executive Committee and board of Directors of Harvard Medical Faculty Physicians at BIDMC, and on the Clinical Operations Executive Committee of BIDMC. The BIDMC Department of Psychiatry is enhanced by the Academic Department of the Massachusetts Mental Health Center (MMHC), the research and academic programs of which have recently been integrated into the BIDMC Department, and programs of which have recently been integrated into the BIDMC Department, and which has an unparalleled reputation in American Psychiatry. The Chairman will be expected to strengthen and expand the existing Departmental clinical programs, to lead academic research in Psychiatry, and to support the continuation of a strong residency program in Psychiatry—the Harvard Longwood Psychiatry Residency Training Program—in collaboration with other area Harvard affiliated hospitals. The applicant should have an M.D., or equivalent degree, be Board Certified in Psychiatry in the US, be a leader in a subspecialty of Psychiatry, and should have a well-developed academic background that will merit appointment as Professor at Harvard Medical School.

Letters of nomination or application outlining experience and career goals, Curriculum vitae, and a list of referees who may be contacted should be sent to:

Albert M. Galaburda, M.D. Chief, Division of Cognitive Neurology Chair, Psychiatry Search Committee Beth Israel Deaconess Medical Center 330 Brookline Avenue, KS-274, Boston, MA 02215 agalabur@bidmc.harvard.edu

Harvard Medical School and Beth Israel Deaconess Medical Center are Equal Opportunity/Africative Action Employers, Women and underrepresented minorities are particularly encouraged to apply.



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DIRECTOR OF PSYCHIATRY



MOUNT SINA SCHOOL OF MEDICINE

Mount Sinai School of Medicine's affiliation with Queens Hospital Center is seeking a Director of Psychiatry to oversee the full range of clinical and administrative functions in the Department of Psychiatry at Queens Hospital Center.

The Department of Psychiatry provides a comprehensive array of clinical services to a vibrant, multicultural community, including a Comprehensive Psychiatric Emergency Program, Inpatient and Outpatient Mental Health Services and Chemical Dependency Services.

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Jasmin Moshirpur, MD
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Queens Hospital Center
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Jamaica, NY 11432
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Chairman Department of Psychiatry

MetroWest Medical Center is seeking a dynamic, creative psychiatrist to lead the Department of Psychiatry. The Chairman will be responsible for providing clinical and administrative leadership to an active department that includes 28 psychiatrists practicing in a 469-bed, two-hospital system, located in the Western suburbs of Boston. The ideal candidate will possess:

- · A strong, demonstrated track record in Physician leadership
- · Board certification in psychiatry
- · Current experience in clinical practice with exceptional clinical skills and commitment to quality
- · Administrative experience in a hospital setting
- · Ability to foster collaborative relationships with local physicians
- · Vision to develop outstanding inpatient and outpatient psychiatric services

The Department of Psychiatry provides a broad range of psychiatric services in a community setting. The hospital has 48 beds devoted to Child, Adult, and Geriatric inpatient care; Evaluation and Referral Services; ECT Services; and a Partial Hospital Program.

MetroWest Medical Center is a fiscally strong medical system that is highly regarded for its commitment to psychiatric care. The MetroWest region of Massachusetts is an attractive and growing area with excellent school systems and convenient access to Boston.

Interested Candidates should send their curriculum vitae in confidence to:

Rebecca Woods, Market VP · Physician Recruitment and Retention Vanguard Health Systems/MetroWest Medical Center 132 Turnpike Road, Suite 200 · Southborough, MA 01772

Inquiries: 508-363-9921 · E-mail: tmcadams@vhsnewengland.com

Psychiatrists - Wisconsin

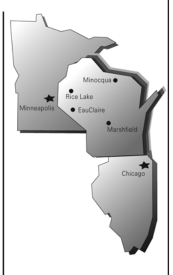
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 St. John's Clinic
 Phone: 800-218-5079
 Fax: 888-290-8300
 E-mail: JAOliver@mercy.net
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Chair of the Department of Psychiatry and Behavioral Sciences Duke University School of Medicine

Duke University School of Medicine invites applications and nominations for the Chair of Psychiatry and Behavioral Sciences. Candidates should have a strong background in academic medicine, including a track record of excellence in patient care, medical education, research and significant leadership experience with peers, trainees and students.

The Department consists of over 400 clinical and research faculty, including psychiatrists, psychologists, medical sociologists, social workers and substance abuse counselors. Clinical services are provided across the entire spectrum of mental illness. Training programs educate medical students, residents, fellows and psychology interns. The Department has over 120 million dollars per year in research support and ranks 4th in funding from the National Institute of Health. Research spans broad areas of social, psychological and biological disciplines focused in behavioral medicine, neurobiology, nicotine and toxicity, epidemiology and intervention trials.

Duke University School of Medicine is among the top research medical schools in the country, annually ranked among the top five schools in NIH funding and among the top 10 by *U.S. News and World Report*. The Duke University Health System's signature clinical facility, Duke University Hospital, is a Magnet hospital located on the Duke Campus and is currently ranked 7th by *U.S. News and World Report*.

The successful candidate should have an M.D. or M.D., Ph.D., academic credentials that qualify for appointment at the rank of Professor with tenure, board certification in psychiatry and eligibility for a license to practice medicine in North Carolina.

Interested individuals should submit a statement of interest and curriculum vitae to: Christopher O'Connor, M.D., Professor of Medicine, Box 3356, Duke University Medical Center, Durham, NC 27710 or via email to: oconn002@mc.duke.edu.



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ALEXANDRIA Strong Clinical Skills. Prefer experience in General Outpatient, Inpatient Psychiatry, and Substance Abuse. CV/Application to heather.ball@va.gov or mail to Heather Ball/Psychiatry Service (116), P.O. Box 69004, Alexandria, LA 71306-9004. For additional questions, please call (318) 466-2958.

SHREVEPORT Prefer experience in Substance Abuse, PTSD. Contact Kathy Arroyo at (318)990-5154 or email at Kathy.arroyo@ va.gov. Email or mail your CV to VAMC, HRMS (05) KA, 510 E. Stoner Ae, Shreveport, LA 71101.

FAYETTEVILLE, FORT SMITH, ARKANSAS; BRANSON, MISSOURI Contact Betty Gray (479)443-4301 ext 5188 or email: betty.gray@va.gov.

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JACKSON, MISSISSIPPI Prefer experience in general psychiatry, including inpatient, outpatient, consultative, or telemedicine psychiatry. Interested candidates should submit a CV to Felicia Owens, Human Resources (05P), VA Medical Center, 1500 E. Woodrow Wilson Dr., Jackson, MS 39216 or Felicia.ovens@va.gov phone: 601-364-1575.

Rush Medical College/Rush University Medical Center

CHAIR, DEPARTMENT OF PSYCHIATRY

Rush Medical College at Rush University Medical Center is seeking outstanding candidates for the position of Endowed Professor and Chair of the Department of Psychiatry. The Department is founded on a long tradition of excellence in clinical care, research and teaching. Currently, there are five endowed chairs in the department. The Department Chair has oversight over the Sections of Adult and Child Psychiatry. Rush is committed to providing outpatient and inpatient psychiatry services, with separate inpatient units for geriatric psychiatry; adult affective disorders, general psychiatry, and child psychiatry. The Chair also has responsibility for fully accredited training programs in Adult Psychiatry and Child Psychiatry.

Candidates must have an outstanding record of commitment to clinical service and research, and substantial administrative experience with an established national reputation as an academic leader. A commitment to advancement of the Department's research mission is also important. In addition, candidates must possess a commitment to innovation in the field and the leadership skills necessary for faculty development and advancement of clinical and academic missions.

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Julie Karstrand
Staff Support for Search/Office of the Dean
Rush University Medical Center
600 South Paulina · Chicago, Illinois 60612

Or preferably electronically to: Julie_Karstrand@rush.edu



CVs should be submitted no later than April 30, 2009

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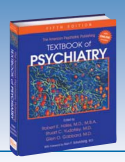
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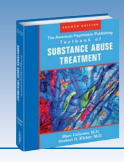
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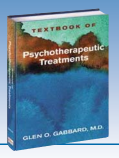
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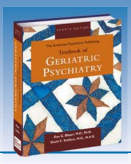
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