LUVOX® (fluvoxamine molecte) 25 mg TABLETS, 50 mg and 100 mg SCORED TABLETS Brief Summary (For full Prescribing Information and Patient Information, refer to package insert.)

INDICATIONS AND USAGE

LIVIQNS Tablets are indicated for the treatment of obsessions and compulsions in adults and children and adolescents (ages 8-17) with Obsessive Compulsive Disorder (OCD), as defined in the DSAHI-R.

CONTRAINDICATIONS

ion of terfenodine, asternizole, cisopride, or pimozide with LUVOX® Tablets is contraindicated (see WARNINGS and PRECAUTIONS). LUVOX® Tablets are contraindicated in patients with a history of hypersensitivity to fluvoxamine maleate.

In patients receiving another serotonin reuptake inhibitor drug in combination with monoamine oxidase inhibitors (MAOI), there have been reports of serious, sometimes fatal, reactions. Some cases presented with features resembling neurolepit medignant syndrome. Therefore, it is recommended that LUVON' Tablets not be used in combination with of MAOI, or within 14 days of discontinuing treatment with a MAOI. After stopping LUVOX° Tablets, or least 2 weeks should be allowed before

14 days of disconnump reunnerment and a starting a MAOI.

Terfenadine, astemizole, disapride, and pimozide are all metabolized by the cytochrome P450IIIA4 isozyme. Increased plasma concentrations of terfenadine, astemizole, disapride, and pimozide cause OI prolongation and have been associated with torsades de pointer-type ventricular tachycardia, sometimes starta. Although it has not been definitively demostrated that fluvoxamine is a potent IIIA4 inhibitor, it is likely to be. Consequently, it is recommended that fluvoxamine not be used in combination with either terfenadine, astemizole, cisapride, and pimozide.

Recommended

Termination

Termination

Description

**Descriptio

combination with either tertenedine, astemizole, cisapride, and pimozide.

Other Potentially Important Drug Interactions. (Also see PREAUTIONS - Drug Interactions). Benzodiazepines: Benzodiazepines melabolized by hapotic audiotina (e.g., alprazolam; midrazolam, inicatolam, etc.) should be used with coution because the clearance of these drugs is ikely to be reduced by fluvoramine. The clearance of benzodiazepines metabolized by gluvoramicinion (e.g., lorazeporm, ovazeporm, ternizeporm) is unlikely to be offected by fluvoramine. Alprazolam: When fluvoramine moleone (100 mg qd) and alprazolam (1 mg qid) were co-administeed to steady, state, plasma concentrations and other pharmacokinetic parameters (AUC, C.m., T.) of alprazolam were approximately hivis those observed when alprazolam was administeed alone; and clearance was reduced by about 50%. The elevated plasma alprazolam concentrations resulted in decreased psychomotor performance and memory. This interaction, which has not been investigated using higher doses of fluvoramine, may be more pronounced in 300 mg daily dose is co-administered, particularly since fluvoramine exhibits non-linear pharmacokinetics over the dosage range 100-300 mg, if alprazolam is and administered with 1UVOX** floates, and decreased psycholam be a condiminated of the contraction of the contracti edministered with LIVDX* Toklets, the initial oliprozolem dosage should be at least behead and itrution to the lowest affective dose is recommended. No dosage adjustment is required for LIVDX** Toklets. Diazepams: The condiministration of LIVDX** Toklets and diazepam is entered to the discrete of both diazepam and its active metabolite. Helementylaterpoor, there is a strong likelihood of substantial occumidation of both species during charactic condiministration. Evidence supporting the conclusion that it is inodivisable to condiministration does not long and diazepam is derived from a study in which healthy volunteers toking 150 mg/day of throwamine were ordinaistered a single or dose of 10 mg of appear. In these subjects (N=8), the clearance of diazepam was reduced by 65% and that of N-desmethyldiazepam to a level that was too low to measure over the coarse of the 2 week long study. It is likely that this experience significantly undescribance may even be more pronounced when it is administrated to higher closers. Accordingly, diazepam and fluvoramine should not ordinaity be condiministration. Amerover, as noted with alprazialam, the effect of fluvoramine may even be more pronounced when it is administrated to higher discrete should be mortificed. The publishment of a single dose of theephylline of the publishment of a single dose of theephylline of the publishment of a single dose of theephylline of single dose of the especial form of the single dose of the usual deliy maintenance dose and plasma concentrations of theophylline is should be mortificed. When they are simple mediate (50 mg) and you are divinished to concentrations of the object like and of the single deliy maintenance dose and plasma concentrations of theophylline is white divoramine mediate (50 mg) and you are officially with verafrain for tow dose, wordning the plasma concentrations of the object like and with fluvoramine mediate. (50 mg) with your dominished concentrations of the object like and were a single part of the single part with ver required for LIVOX** Toblets. *Warfarins*. When flivoxorania maleate 5.00 mg trol was administered concomitantly with verdorin for bow exeks, worfarin plasma concentrations increased by 98% and prothrombin firms were prolonged. Thus parients receiving and anticoagulants and LIVOX** Toblets should have their prothrombin firms manitared and their anticoagulant dose adjusted accordingly. No desage adjustment is required for LIVOX** Toblets.

PRECAUTIONS

General

General
Activation of Mania/Hypomania: During premarketing studies involving primarily depressed potients, hypomania or mania occurred in approximately
1% of potients treated with fulveractions. Activation of mania/hypomania has also been reported in a small proportion of patients with major affective
disorder who were treated with other marketed antidepressons. As with all antidepressons, LIVOX* Tables should be used cauthously in patients with the history of seazures. It should be discontinued in any patient who develops seizures. Suicide: The possibility of a saided extension
is inherent in patients with depressive symptoms, whether these occur in primary depression or in association with another primary disorder such as O.D.
Closs supervision of high risk patients should accompany initial dug therapy. Prescriptions for UNOX* Tables should be written for the smallest quantity of
tablest consistent with good patient management in order to reduce the risk of overdose. Use in Patients with Committain Illness: Committed committed expension with UNOX* Tables is patient with systemical less is initial. Custom is obvised in deministrating IUVOX** Tables
to patients with diseases or conditions that could offert hemodynomic responses or metabolism. LIVOX** Tables have not been evaluated or used to one
participated in proteins with a potential studies during the product's premarketing testing. Evolution of the electrocordiograms for petients with depression or OOD who
participated in premarketing studies revealed no differences between thoursamine and placeton in the response to proteins in patients with his
dysfunction, fluvocaminia clearrance was decreased by approximately 30% LIVOX** Tables: Interference
harmanian for Patients: Proteins of the proteins

Information for Patients: Physicians are advised to discuss the following issues with notions for whom they prescribe LEVOX® Tablets: Interference Internation for rements: ritigation are underly document of sources are inversing assess with parameter or more may presume curve. In most, an extraction with Cognitive or Motor Performance: Since any sychocotive drug may import indigenent, thinking, or motor skills, pointent be causined about operating hazardous machinery, including outernobiles, until they are certain that LUVOX** Tablest therapy does not odversely affect their ability to acoust opening nucroscon interating; incoming autonouses, unin mey use certain that DVDV." Tables metaly does not observes direct that consequently engaged in the configuration of the consequently and the consequently of the consequently interest the consequently interest the consequently of the consequen

Laboratory Tests: There are no specific laboratory tests recommended

Drug Interactions: There have been rare postmarketing reports describing patients with weakness, hyperreflexic, and incoordination following the use of a selective serotonin reuptake inhibitor (SSRI) and sumatriplan. If concomitant treatment with sumatriplan and SSRI (e.g., fluoxetine, fluoxeanine, of a selective seritorian recipide imbatrial (SSAI) and summitted in It concomitral treatment with summittation and SSAI (e.g., Buozetine, Buvozeniae) programseine, seritorials is chaircally community of propriet in deviced. Protential interactions with drugs that inhibit or are Metabolized by Cytochrome P450 Isozymes: Based on a finding of substantial interactions of fluvozamine with certain drugs and limited in vitro data for the IIIA4 Sozyme, it appears that fluvozamine interactions is possible with drugs having a narrow therapeutic ratio such as serienciae, estimated, estignitie, or prinaride, warfalin, theaphylline, certain benzediarepines and phenytria. If IUVOX* liabels are to be administered together with a drug that is eliminoted via oxidative metabolism and has a narrow therapeutic window, pasma levels and/or pharmacodynamic effects of the latter drug. a audy ain a enimanen via obtainive meruculari and rus a initiari wherepourit window, pastral eves and /or pinarmacopyramic energs of the interest as should be maintained classify, it least mild steady-state conditions are reached. Pleas see compiler perscribing information for recommendations regarding CNS drugs such as monoarnine oxidase inhibitors, alprazolam, diazepam, lorazepam, lithium, trystopham, clazapine, alcohol, tricyclic antidepressants, carbamacepine, methodors, and other drugs with as theosphiline, proparoidal and other beta-diockers, variorini, digoxin, difficzem. Effects of Smaking on Flavoramine Metabolisms: Smokes had a 25% increase in the metabolism of throwsamine componed to nonsmakers. Electroconvulsive Therapy (ECT): There are no clinical studies establishing the benefits at risks of combined use of ECT and flavoxamine maleate.

Therefore (ECLT): There are no clinical sholes strobishing the benefits or risks of combined use of ECL and fluvoxamine maleate. Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenesis: There is no evidence of corcinogeneity, mutagenicity or impairment of fertility with fluvoxamine maleate. There was no evidence of carcinogenesis: There is no evidence of corcinogeneity in rust headed orably with fluvoxamine maleate for 30 months or homostes headed orably with fluvoxamine maleate for 30 months or homostes headed orably with fluvoxamine maleate for 30 months or homostes headed orably with fluvoxamine maleate for 30 months or homostes headed orably with fluvoxamine maleate for 30 mg/kg in comparison or the study from a minimum of 160 mg/kg to a maximum of 1240 mg/kg in homostes. The maximum house of 240 mg/kg is approximately 6 limes the maximum human daily dose on a mg/m² basis. Mutagenesis: No evidence of mutagenic potential was observed in a more of Fertility: In entility studies of made and female rats, up to 80 mg/kg/doy orally of fluvoxamine maleate (approximately 2 times the maximum human daily dose on a mg/m² basis) had no effect on mating performance, dworfion of gestation, or pregnancy rate.

Pregnancy
Tevrologenik Effects: Pregnancy Category C: In teutology studies in ruts and robbits, daily and doses of fluoroxamine maleate of up to 80 and 40 mg/kg, respectively (approximately 2 times the maximum human daily dose on a mg/m² basis) caused no fetal inafformations. However, in other reproduction studies in which pregnant rats were dosed through wearing there was (1) an increase in purp mortality or birth (seen at 80 mg/kg and dose) but not at 20 mg/kg, and cally observes in postability but weights (seen at 160 but not at 80 mg/kg) and (20) decreases in postability weights (seen at 160 but not at 80 mg/kg) and (seen at all doses, lowest dose tested = 5 mg/kg). (loses of 5, 20, 80, and 160 mg/kg are approximately 0.1, 0.5, 2, and 4 times the maximum human daily dose on a mg/m² basis.) White the results of a cross-fostering study implied that at least some of these results likely occurred secondarily to maternal basis, to, the relief of any feet some or purps could not be ruled out. These are no adequate and well-nothfalled studies in pregnant women. Fluoroximine maleate should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Labor and Delivery: The effect of fluvoxamine on labor and delivery in humans is unknown

Nursing Mothers: As for many other drugs, fluvoxomine is secreted in human breast milk. The decision of whether to discontinue nursing or to discontinue the drug should take into account the potential for serious odverse effects from exposure to fluvoxomine in the nursing infant as well as the potential benefits of LUVOX* (fluvoxomine maleate) Toblets therapy to the mather.

Pediatric Use: The efficacy of fluvoxamine molecte for the treatment of Obsessive Compulsive Disorder was demonstrated in a 10-week multicenter placebo controlled study with 120 outpatients ages 8-17. The adverse event profile observed in that study was generally similar to that observed in adult studies with fluvoxamine (see ADVERSE REACTIONS).

Decreased appetite and weight loss have been observed in association with the use of fluvoxamine as well as other SSRIs. Consequently, regular monitoring of weight and growth is recommended if treatment of a child with an SSRI is to be continued long term.

Geriatric Use: Approximately 230 patients participating in controlled premarketing studies with LUVOX® Tablets were 65 years of age or over. No overall

differences in safety were observed between these patients and younger patients. Other reported clinical experience has not identified differences in response between the elderly and younger potents. However, filtwoomnine has been associated with sevent cases of clinically significant hypometric many of the elderly compared to younger potents, and greater sensitivity of PRECAUTIONS, General). Furthermore, the cleanance of filtwoomnine is decreased by about 50% in elderly compared to younger patients, and greater sensitivity of some older individuals also cannot be ruled out. Consequently, LUVOX® Tablets should be slowly titrated during initiation of the

ADVERSE REACTIONS

Associated with Discontinuation of Treatment: Of the 1087 OCD and depressed patients treated with fluvoxamine maleate in controlled clinical trials conducted in North America, 22% discontinued treatment due to an adverse event.

Incidence in Controlled Trials - Commonly Observed Adverse Events in Controlled Clinical Trials: LUYOX® Tablets have been studied in controlled thirds of OCI (N=320) and depression (N=1350). In general, adverse event rates were similar in the two data sets as well as in the pediatric.

OCD study. The most commonly observed adverse events associated with the use of LUVOX* Tablets and likely to be drug-related (incidence of 5% or greater and at least twice that for placebo) derived from Table 1 were: somnolence, insomnia, nervousness, tremor, nausea, dyspepsia, narvexia, varning,

and at lost twice that for glarebol derived from Table I were: somoelence, insomalic, nervousness, herence, nausea, dyspessia, nonexia, vanning, abnormal ejecutation, asthenia; and sweeting. In a pool of two studies involving only potients with OCD, the following additional events were identified using the clover rule: dry mouth, decreased libidia, uninary frequency, anagasmia, thinitis and taste preversion. In a study of pediatric patients with OCD, the following additional events were identified using the above rule: agrintion, depression, dysmenanthea, floridence, hyperkinesia, and rash. Adverse Events Occurring at an Incidence of 1965: fable I enumentes adverse events that occurred at a frequency of 1% or more, and were more frequent than in the placebo group, among patients thereted with LUVOX® Tablets in two short-term placebo controlled OCD trials (10 week) and depression trials (6 week) in which patients were dosed in a range of generally 100 to 300 mg/day. This table shows the precentage of patients in early only which and telest one occurrence of an event of some time during their freatment. Reported obserse events were clossified using a standard OCSTART-based Dictionary terminology. The prescriber should be oware that these figures comont be used to predict the incidence of side effects in the course of used medical practice where potient democratistics and other factors may differ from those that prevailed in the clinical his-clinical his-clinical mixed physician with some basis for estimating the relative contribution of drug and non-drug factors to the side-effect incidence rate in the coolation studied.

on provice me prescrioning preyconn with some basis for estimating the reactive continuation of oxig and non-ring factors to me solve-errer indocence rine in peopularion studied.

Table 1: TREATMENT-MERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN ADULT OCD AND DEPRESSION POPULATIONS COMBINED! (fluvocomine (N=892) is, placeto (N=7/8) by patients—percentage): BODY AS WHOLE: Hoodobre (22 vs. 20); Ashenic (14 vs. 6); Flu Syndrome (3 vs. 2); Chills (2 vs. 1); CARDIOVASCULAR: Polpitotions (3 vs. 2); DIGESTIVE SYSTEM: Nausea (40 vs. 14); Borrhea (11 vs. 7); Constipation (10 vs. 8); Dyspepsia (2 vs. 8); Discomina (2 vs. 10); Dyspepsia (2 vs. 9); Depression (2 vs. 10); Abustive (3 vs. 1); Anxiety (5 vs. 3); Viscolitation (2 vs. 10); Bypersian (2 vs. 10); Abustive (3 vs. 10); Depression (1 vs. 6); Flemon (5 vs. 11); Anxiety (5 vs. 3); Viscolitation (3 vs. 10); Dyspepsia (1 vs. 10); Dyspepsia (2 vs. 10); Dyspepsia

were: astheraia, abnormal ejeculation (mostly delayed ejeculation), anxiety, infection, thintis, anagasmia (in males), depression, hidio decessed, phayangins, gather, impotence, myoclorus/which, thirst, weight loss, leg camps, myolgia and urinary retention. These events are listed in order of decreasing rates in the OCD thirds.

Other Adverse Events in OCD Pediatric Population: In Pediatric patients (N=57) neaded with LUVOX[®] Tablets, the overall profile of adverse events is similar to that seen in adult studies. Other reactions which have been reported in two or more pediatric patients, and were more frequent than in the placebo group were: abnormal thinking, cough increase, dysmenarther, endlymosis, emotional fability, epistaxis, hyperkinesia, infection, manic reaction, rash; sinusitis, and

Vital Sign Changes: Comparisons of fluvoxamine maleate and placebo groups in separate pools of short-term OCD and depression trials on (1) median n boseline an various vital signs variables and on (2) incidence of potients meeting criteria for potentially important changes from base various vital signs variables revealed na important differences between fluvoxamine maleate and alaceba.

Laboratory Changes: Comprisons of Photosomine melecte and placebo groups in separate pools of short term OCD and depression trials on (1) median change from boseline on various serum chemistry, hemotology, and uninolysis variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various serum chemistry, hematology, and urinalysis variables revealed no important differences between fluvoxami

malaste and placebo.

ECG Changes: Comparisons of fluvoxamine maleate and placebo groups in separate pools of short-term OCD and depression trials on {1} mean change from baseline on warious ECG variables and on {2} incidence of patients meeting criteria for potentially important changes from baseline on various ECG variables revealed no important differences between fluvoxamine maleate and placebo.

Other Events Observed During the Premarketing Evaluation of LUVOX* Tablets: During premarketing clinical trials conducted in North America and Europe, multiple dosses of fluvoxamine melaste were administered for a combined total of 2737 patient exposures in patients suffering OCD or Najar Depressive Boardes. Untoward ovents associated with this exposure were recorded by clinical investigators using descriptive terminology of their pown choosing. Consequently, it is no possible to provide a meninegial existent of the popontion of individuode seprendering adverse without first grouping similar types of untoward events into a limited (i.e., reduced) number of standard event categories. In the tabulations which follow, a standard COSIAR/based Dictinary terminology has been used to classify reported adverse events. If the COSIAR/term for an event was so general as to be uniformative, if was reproced with no more informative term. The frequencies presented, thereofore, perseentth proception of the \$2.25 and \$2.05 COSINATORED LINCOLOGY Intellined by an extent beat or classify; ejected unweste events. If the COSINATA feath of an event was 50 generals as to uninformative, if was repicted with a more informative term. The frequencies presented, therefore, represent the proportion of the 2737 points apposure to multiple doses of fluvocomine molecule who experienced on event of the type cited on of least one occasion while receiving fluvocomine molecule reported events or included in the list below, with the following exceptions: 1) those events bearing fished in Table 1, which thoulates incidence rates of common odverse experiences in placebo-controlled OCD and depression clinical thick, one excluded; 2) those events for which a drug coose was considered remate (i.e., neoplasia, gastrointestinal carainome, herpes simplex, herpes zoster, application site reaction, and unintended pregnancy) are armitted; and 31 events which were reported in only one patient and judged to not be potentially serious are not included. It is important to emphasize that, although the events reported did occur during treatment with fluvoxamine maleate, a causal relationship to fluvoxamine maleate has not been established. Events are events reported did o'ccur during heatment with fluvoxamine maleate, a causal relationship to fluvoxamine maleate has not been established. Events are further dossified within body system categories and enumented in order of decreasing frequency using he following definitions: frequent otherses were to effer one or more accussions in orl least 1/1000 potients; frequent oderse events are these accurring breween 1/100 and 1/1000 potients, and rure adverse events are those occurring on in less than 1/1000 potients. Body as a Whale: Frequent: accidental injury, maloise; infrequent: allergic reaction, nuck pain, nuck rigidity, overdose, photorestifity proctron, suicide attempt, Rure: cyst. pelvic pain, sudden death. Cardiovascular Systems: Frequent: hypertension, typocresion, syropoe, todaycardia; infrequent: adiapre, petrits, bodycardia, indervous control reports, bodycardia; indervous accordance systems: Frequent: charges petrits, bodycardia, place frequent cardiovascular disease, coid extremities, conduction delay, heart failure, myocardial inforction, pollor, pulse irregular, SI segment changes; Rure: Al block, curebrowscular ocident, corrowary terity desse, embolus, pericardiis, pillebitis, pulmonary inforction, suproventificate extracysioles. Digestive Systems: Frequent: Reveited liver transmisses; Infrequent: chings, ructrion, espolitagilis, gastriss, postmentering administrational destruction, pundice. Endocrine System: Infrequent: hypothyroidism; Rure: glotte, Henric and Lymphatic Systems: Infrequent: development, purplus, Metabolic and Murritional Systems: Infrequent: development, purplus, Metabolic and Murritional Systems: Frequent: development, purplus, Metabolic and Murritional Systems: Frequent, polycolomin, hypotherina, leadinous controlute, tenorynoitis; Rure orthoosis, mypothyroidism; Rure: dollares mellins, hypoglycemia, hypoglycemia, hypoglycemia, pulse, controlute, lenorynoitis; Rure: orthoosis, mypodlym, pulhologia fincture. Nervous System: Frequent annessa, apothy. hypoglycemia, hypokolemia, loctate dehydrogenose increased. **Mosculoskeletal System:** Infraquent: arthrolgia, arthritis, bursitis, generalized muscle sporan, myrstheria, tendinous centrocture, tenosynovitis, Rare criticosis, mypodinis, pathological fracture. **Moreous System:** Frequent: armasia, generalized muscle sporan, myrstheria, tendinous centrocture, tenosynovitis, Rare criticosis, myrodinis, pathological fracture. **Moreous System:** Frequent: armasia, and deliaria, delatisian, debescandization, drug dependence, dyskinesia, dystemia, emotipanal balbity, euphoina, extrugromadal syndrome, agait unsteady, hollucinations, hemiplegia, hossilisty, hypersaminia, hypochordriacis, hyperoina, hysteria, incoadrantion, increased silvertion, increased blido, neuraligia, paralysis, paramoid reaction, phobia, psychosis, sleep disorder, suport, britching, vertigo; Rare: dainesia, cama, fibrillations, mutism, dossissions, reflexes decreased, starred speech, tardive dyskinesia, torticulis, trismus, withdrawal syndrome. **Respiratory System:** Frequent: Cough increased, sinustis, Infraquent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, playedent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, playedent: acre, alopeia, dry skin, ezerno, exfloative dermathis, fruunculosis, seborthes, skin discolarian, utticunis. **Special Senses:** Infraquent: acre, properties, skin, acre, playedent: acre, alopeia, dry syes, experimentation acre Based on the number of females. 2Based on the number of males.

Non-US Postmarketing Reports: Voluntary reports of adverse events in patients taking LUVOX* Tablets that have been received since market introduction and are of unknown causal relationship to LUYOX® Tablets use include: toxic epidermal necrolysis, Stevens-Johnson syndrome, Henoch-Schoenlein purpura, bullous eruption, priopism, agranulocytosis, neuropothy, aplastic anemia, anaphylactic reaction, hyponatemia, oxute renal failure, henatitis, and severe akinesia with fever when fluvoxomine was co-administered with antiosychotic medication

OVERDOSAGE

Refer to package insert (15E Rev 5/99) for overdosage information.

DOSAGE AND ADMINISTRATION

Refer to package insert (15E Rev 5/99) for dosage and administration information.

Solvay Pharmaceuticals Marietta, GA 30062 Rev 6/99 (1280/1285 15F Rev 5/99)

Solvay **Pharmaceuticals**

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"My doctor diagnosed obsessions and compulsions and prescribed LUVOX® Tablets."



- ▼ IMPROVES OBSESSIVE-COMPULSIVE SYMPTOMS IN ADULTS, CHILDREN, AND ADOLESCENTS^{2,3}
- ▼ LOW INCIDENCE OF SEXUAL DYSFUNCTION IN ADULTS⁴
 LUVOX® Tablets vs placebo: decreased libido 2% vs 1%; delayed ejaculation 8% vs 1%; impotence 2% vs 1%
- ▼ LOW INCIDENCE OF AGITATION IN ADULTS⁴ 2% vs 1% for placebo

In adults, the most commonly observed adverse events compared to placebo were somnolence 22% vs 8%; insomnia 21% vs 10%; nervousness 12% vs 5%; nausea 40% vs 14%; asthenia 14% vs 6%⁴

In children and adolescents, the most commonly observed adverse events compared to placebo were: agitation 12% vs 3%; hyperkinesia 12% vs 3%; depression 5% vs 0%; dysmenorrhea 7% vs 3%; flatulence 5% vs 0%; rash 7% vs 3%⁴

Concomitant use of LUVOX® Tablets and monoamine oxidase inhibitors is not recommended.4

Fluvoxamine should not be used in combination with terfenadine, astemizole, cisapride, or pimozide.4

As any psychoactive drug may impair judgment, thinking, or motor skills, patients on LUVOX® Tablets should be advised to exercise caution until they have adapted to therapy.4

References: 1. Physician Drug & Diagnosis Audit (PDDA) and Source™ Prescription Audit (SPA) August 1999-September 1999. Scott-Levin, a division of Scott-Levin PMSI Inc. 2. Goodman WK, Kozak MJ, Liebowitz M, et al. Treatment of obsessive-compulsive disorder with fluvoxamine: a multicentre, double-blind, placebo-controlled trial. *Int Clin Psychopharmacol*. 1996;11:21-29. 3. Data on file, Study in Children and Adolescents (Report No. CR200.0116), Solvay Pharmaceuticals. 4. LUVOX® Tablets Full Prescribing Information.

VISIT OUR OCD WEB SITE AT www.ocdresource.com

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Please see brief summary of prescribing information on adjacent page.

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First-line SSRI therapy for obsessions and compulsions