LUVOX [®](fluvoxamine maleate) Tablets

Brief Summary (For full Prescribing Information refer to package insert.)

INDICATIONS AND USAGE

LUVOX Tobles are indicated for the nearment of classisons and compulsions in patients with Obsessive Compulsive Disorder (OCD), as defined in the DSMHR. The abses-sions or computations cause marked datess, are inter-consuming, or significantly interfere with social or accupational functioning. The efficacy of LUVOX Toblets was estab-lished in two To-week triads with obsessive computative compactive swith the diagnosis of Obsessive Computative Disorder as defined in DSMHR. Obsessive Computative Disorder as defined in DSMHR. parter in the 10 hear nas mental calibration strategies and position in the magnetic strategies and the stra usefulness of the drug for the individual patient.

CONTRAINDICATIONS

Co-administration of terfenadine, asternizale, or cisapride with LUVOX Tablets is contraindicated (see WARNINGS and PRECAUTIONS). LUVOX Tablets are contraindicated in with a history of hypers WARNINGS

potents with a histry of hypersensitivity to fluvocamine modelle. WARNINGS Potential for interaction with Monocamine Oxidese Inhibitors. In patients receiving another sorotomin receptate inhibitor drug in combination with menocamine oxides inhibitors (MOD), there have been reports of serious, sametimes fatal, reactions including hyperthemia, rightity, myodoway, nationanic instability with possible rapid fluctuations of vital signs, and mental status changes that include extense ogitation pro-gressing to deliver and non. These reactions have also been reported in patients who have discontinued that drug and have been started on a MAOI. Some cases presented with features resonabling neurologic in includies, and mental status changes that include extense ogitation pro-gressing to deliver allowed before or writing in 14 days of discontinuing treatments with a MAOI. After stopping LUVOX Tablets, at least 2 weeks shade to allowed before starting of MAOI. Potential Verfandine, Astenzize, and Changeride Interactions. Torfanodine, astenizade, and cisquicite are all methodized by the cytochrome PS501IMA (stopyme, and it has been demonstrated that ketocourcile, a potent inhibitor of IIMA, blocks the methodism of these drugs, machine and exacutivities of parent drug. Increased pleama concentrations of terfenodine, astenizade, and cisquicide case of prolongation and lossabels de pointer-type vanitoker tady and that is know, a substantial pharmachileritic interactions in a been definitively demonstrated that flavocanies is potent IIMA inhibitor, is is likely to be, given the substantial interactions of Abought has not been definitively demonstrated that flavocanies is potent IIMA inhibitor, is is likely to be, given the substantial interactions of has and precasions. Consequently, its recommended that flavocanies. The demance demanced interactions of the yournation of the same device (AUTIONS) and PRECAUTIONS). Other Potentially important Drug Interactions (do see reflectively table, pice induces by howar

Learners are a new process parame concernments are prommoscience pormeters SUL, Cytoge, Tyr.) of objacation were opportunitely twice those indexended when advacation were solutinisted alone; and clearance was tacked by about 50%. The elevated phasma diparadiam concentrations resulted in decreased psychomator perfor-mance and memory. This interaction, which has not been investigated using high does of fluoraname, may be mare parameters the subscription of the su

PRECAUTIONS

for LUXO Tables. **PRECAUTIONS General Activities of Mania/Hypomonic**: During premarketing studies involving primarily depressed patients, hypomania or mania accurred in approximately 1% of patients treated with housemine. Activation of mania/hypomania has data been reported in a small proportion of patients with india of field the disade who were treated with their marketed mice presents. Subject the intervention of mania/hypomania has data been reported in a small proportion of patients with india of field the disade who were treated with their marketed mice presents in 0.2% of housemane treated presents. UNOX folders should be used controlsy in patients with a history of mania. Subject moves the treated with their sciences were created in 0.2% of housemane treated presents. UNOX folders should be used controlsy in patients with a disease-serve presents in 0.2% of house the sciences in the present in 0.2% of house a science in the science i

There are no specific laboratory tests recommended.

Laboratory Fast There are a costefic blockney less terrormendel. Traj liveractions Protectial interactions with Drags that habit or are Metabolized by Cytochrone P450 Iszaynes: Multiple hepatic cytochrone P450 ((YP450) exymes are involved in the individue blocknession of a large number of structurely different drugs and indepances compounds. The evaluable knowledge concerning the relationship of fluxoannine and the CYP450 enzyme system has been abtained mostly from phormacokinet intercofon states could be involved in the term of the social of the

the maximum human daily dose on a mg/m² basis. **Matagenesis:** No evidence of mutagency potential was observed in a mouse microaudeus test, an in who dra-masone obsenation test, or the Ames microbial mutagen test with an writout metadolic activation. **Impairment of Fortility:** In fenility studies of male and fenale rats, up to 80 mg/tgg/day cally of lluvacamine maleate, (opproximately 2 times the maximum human daily dose on a mg/m² basis) had no effect on maning performance. duration of gestation, or pregnancy rate.

duration of epsithinor, or programicy retre. **Prespinacy Prespinacy Prespinacy**

The effect of Buyoyamine on Jabor and delivery in humans is unknown

Nursing Mothers

As for many other drugs, fluoroamine is secreted in human breast milk. The decision of whether to discontinue nursing or to discontinue the drug should take into account the paternial for serious adverse effects from exposure to fluoroamine in the nursing infant as well as the paternial benefits of LUVOXTM Tablets therapy to the mother. Pediatric Use

Safety and effectiveness of LUVOX Tablets in individuals below 18 years of one have not been established

Geriatric Use

Contract Case Approximately 230 patients participating in controlled premarketing studies with LUVOX Tablets were 6.5 years of age or over. No overall differences in safety were observed between these patients and youngar patients. Other reported chincil expensions has not identified differences in response between the elderly and youngar patients. However, the clearance of fluxoamine is decreased by about 50% in elderly compared to younger patients and greater sensitivity of some older individuals also connor be ruled our. Consequently, LUVOX Tables should be solvely theselwh interfation of the target.

ADVERSE REACTIONS

Associated with Discontinuation of Treatment – Of the 1087 O/D and depressed patents neated with Ruoxamine molecule in controlled charical traits conduct-ed in Narth America, 22% discontinued heatment due to an adverse event. The most common events (≥1%) associated with discontinuation and considered to be drug related (i.e., those events associated with drogout at least twice that of placebo) included hendache, asthenia, abdominal pain, nausea, vomitting, diarrhea, dyspessio, anorex-

ear in thin Aneter, 22% ascontrust in terminer due to an overse event. In the most common events, 12% by assocrates with ascontrulation for most events of the dispetision of the backby, related the backby, assocrated with disposal and early executive of the process of the backby end the backby, end th to event rates in OCD and depression studies were, asthenia, abnormal ejaculation (mastly debyed ejaculation), anxiety, infection, rhimits, anargesmia (in males), depres-sion, labida decreased, pharmajnis, againtain, impotence, mycolanus/hiniti, thist, weight loss, leg camps, mylagia and uninary retention. These events are listed in acter of decreasing mars in the OCD trial.

Vocations of fluoreanine molecte and placebo groups in separate pools of short+term OCD and depression trials on (1) median change from baseline on various with signs variables and on (2) incidence of potients meeting criteria for potentially important changes from baseline on various with signs variables revealed no important differences. between fluvoxamine maleate and placebo.

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Laboratory Changes Comparisons of fluvoxamine molecute and placebo groups in separate pools of short-term OCD and depression trais on (1) median change from baseline on various serum chemistry, hematology, and uninalysis variables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various serum chemistry, hematology, and uninalysis variables revealed no important differences between fluvournine maleade and placebo.

ECG Changes

Comparisons of fluvovamine maleate and placebo groups in separate pools of short-term OCD and depression triads on (1) mean change from baseline on various ECG vori ables and on (2) incidence of patients meeting criteria for potentially important changes from baseline on various ECG voriables revealed no important differences between

Table 2: TREATMENT-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COM-

flowcamine molecte and placeb.
Table 2: TREATINET-EMERGENT ADVERSE EVENT INCIDENCE RATES BY BODY SYSTEM IN OCD AND DEPRESSION POPULATIONS COM-BINED¹ (divocamine vs. picetalo by priemtegerandpa): BODY AS VMPOLE: Hostadie (22 vs. 20), kothenio (14 vs. 6); Flo Syndhom (00 vs. 8); Doppadie (10 vs. 5); CARDIOVASCULAR: Polyintors (3 vs. 2): OBGENTVE SYSTEM: Nausee (40 vs. 14); Danhad (14 vs. 6); Flo Syndhom (10 vs. 8); Doppadie (10 vs. 5); CARDIOVASCULAR: Polyintors (3 vs. 2): DBGENTVE SYSTEM: Nausee (40 vs. 14); Danhad (14 vs. 6); Flo Syndhom (20 vs. 12); Shoreita (15 vs. 2); CARDIOVASCULAR: Polyintors (3 vs. 2): DBGENTVE SYSTEM: Nausee (40 vs. 14); Danhad (14 vs. 6); Flow Syndhom (20 vs. 13); Naudiathim² (3 vs. 2); Campania (14 vs. 6); Flow Syndhom (21 vs. 6); Terror (5 vs. 1); Ankivous (5 vs. 3); Vasodiathim² (3 vs. 1); Bepersian (12 vs. 6); Terror (5 vs. 1); Ankivous (5 vs. 3); Vasodiathim² (3 vs. 1); Bepersian (12 vs. 6); Terror (5 vs. 1); Ankivous (5 vs. 3); Vasodiathim² (3 vs. 2); UROSGENTATE: Storense (2 vs. 1); Vasoti (2 vs. 0); SURIM: Swenting (7 vs. 3); SECELAL SENSES: Toste Pavessian (3 vs. 1); Anhiyopis' (3 vs. 2); UROSGENTAL: Abroard Epolation (2 vs. 1); Vasot (2 vs. 0); SURIM: Swenting (7 vs. 3); SECELAL SENSES: Toste Pavessian (3 vs. 1); Anhiyopis' (3 vs. 2); UROSGENTAL: Abroard Epolation* (3 vs. 1); Vasot (2 vs. 0); SURIM: Swenting (7 vs. 3); SECELAL SENSES: Toste Pavessian (3 vs. 1); Anhiyopis' (3 vs. 2); UROSGENTAL: Abroard Epolation* (3 vs. 1); Vianary Freezuency (3 vs. 2); Impotence* (2 vs. 0); Unary Retention (1 vs. 0). Finance on event of the type devia on a field on a clied on a custon while receiving fluxoamine molecule. All reported events are included in the ist belowing exception: 1) those events cliendow tisken in the log with the black with the black of the time to the section on a down sense included. The antibact has a finance the section is the receiving and an interest of pregnancy are united. An all together and the set expended in only one patient and judget to not be potentially series are not included. It is importent to any set of the set of the section is the rectain, and unitered pregnancy are united, and 3) exerts which were reported in only one patient and judget to not be potentially series are not included. It is importent to any set of the section is the rectain and patient to the accessing integration of the area sets blacked. The transmission of the section is the section of the section of the section is the section of the section is the section of the sectin of the section of the se

1 Based on the number of females.; 2 Based on the number of males.

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 Yoluntary reports of adverse events in patients taking LIVOX Tablets that have been received since market introduction and are of unknown causal relationship to LUVOX
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4E1252 Rev 9/95

Reference: 1. Data on file. Solvay Pharmaceuticals. Inc.



Solvay Pharmaceuticals

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ESTABLISHED THERAPY FOR OCD

EFFECTIVE CONTROL OF OBSESSIONS AND COMPULSIONS^{1*}

LOW INCIDENCE OF AGITATION

 $(2\% vs 1\% \text{ for placebo})^1$

LOW INCIDENCE OF SEXUAL DYSFUNCTION

LUVOX® Tablets vs placebo[†]: decreased libido 2% vs 1%; delayed ejaculation 8% vs 1%; anorgasmia 2% vs 0%; impotence 2% vs 1%

FAVORABLE SAFETY PROFILE

- Relatively low incidence of anticholinergic side effects in controlled trials of OCD and depression, LUVOX[®] Tablets *vs* placebo¹: dizziness 11% *vs* 6%; constipation 10% *vs* 8%; dry mouth 14% *vs* 10%
- 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%, abnormal ejaculation 8% vs 1%, asthenia 14% vs 6%¹
- Concomitant use of LUVOX[®] Tablets and monoamine oxidase inhibitors is not recommended¹

FLEXIBLE DOSING

Initial Dose: 50 mg once a day HS Dose Range: 100 to 300 mg/day

COMPREHENSIVE SAFETY DATABASE

(Worldwide Exposure for Reporting Overdose[‡])¹

- Data from 40 countries
- Over 12 million patients treated
- More than 37,000 patients studied in clinical trials

IUVOXAMINE MALEATE SOME & 100 Mg fluvoxamine maleate SOME & 100 Mg some D TABLETS A SELECTIVE SEROTONIN REUPTAKE INHIBITOR

> *Effectiveness not established beyond 10 weeks in controlled trials. [↑]Parameters occurring ≥ 1% with fluvoxamine maleate. [‡]Prescribers should write the smallest tablet quantity consistent with good patient management to reduce overdose risk. *Please see brief summary of prescribing information on adjacent page.*

OUBTS

LEAN

FEARS

WORRY