Lustral® Tablets 50mg
Lustral® Tablets 100mg

WARNING: SUICIDAL THOUGHTS AND BEHAVIORS
Antidepressants increased the risk of suicidal thoughts and behavior in children, adolescents, and young adults in short-term studies. These studies did not show an increase in the risk of suicidal thoughts and behavior with antidepressant use in patients over age 25; there was a reduction in risk with antidepressant use in patients aged 65 and older [see Warnings and Precautions (4.4)].

In patients of all ages who are started on antidepressant therapy monitor closely for clinical worsening and emergence of suicidal thoughts and behaviors. Advise families and caregivers of the need for close observation and communication with the prescriber [see Warnings and Precautions (4.4)].

1. NAME OF THE MEDICINAL PRODUCT

Lustral® tablets 50 mg

Lustral® tablets 100 mg

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film-coated tablet contains sertraline hydrochloride equivalent to 50 mg or 100mg sertraline.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet

Sertraline 50 mg film-coated tablets are white, capsular shaped, film-coated scored tablets marked ‘ZLT 50’ on one side and ‘PFIZER’ on the other. The tablet can be divided into equal doses.

Sertraline 100 mg film-coated tablets are white, capsular shaped, film-coated tablets marked 'ZLT 100' on one side and 'PFIZER' on the other.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications
Sertraline is indicated for the treatment of symptoms of depression in patients with or without a history of mania. Following satisfactory response, continuation with sertraline therapy is effective in prevention relapse of the initial episode of depression or recurrence of further depressive episodes.

4.2 Posology and method of administration

Posology

*Initial treatment*

Sertraline treatment should be started at a dose of 50 mg/day.

*Titration*

Patients not responding to a 50 mg dose may benefit from dose increases. Dose changes should be made in steps of 50 mg at intervals of at least one week, up to a maximum of 200 mg/day. Changes in dose should not be made more frequently than once per week given the 24-hour elimination half life of sertraline.

The onset of therapeutic effect may be seen within 7 days. However, longer periods are usually necessary to demonstrate therapeutic response.

*Maintenance*

Dosage during long-term therapy should be kept at the lowest effective level, with subsequent adjustment depending on therapeutic response.

Longer-term treatment may also be appropriate for prevention of recurrence of major depressive episodes (MDE). In most of the cases, the recommended dose in prevention of recurrence of MDE is the same as the one used during current episode. Patients with depression should be treated for a sufficient period of time of at least 6 months to ensure they are free from symptoms.

*Paediatric patients*

Efficacy is not shown in paediatric major depressive disorder.

*Use in elderly*

Elderly should be dosed carefully, as elderly may be more at risk for hyponatraemia (see section 4.4).

*Use in hepatic insufficiency*

The use of sertraline in patients with hepatic disease should be approached with caution. A lower or less frequent dose should be used in patients with hepatic impairment (see section 4.4). Sertraline should not be used in cases of severe hepatic impairment as no clinical data are available (see section 4.4).

*Use in renal insufficiency*
No dosage adjustment is necessary in patients with renal insufficiency (see section 4.4).

Method of administration
Sertraline should be administered once daily, either in the morning or evening. Sertraline tablet can be administered with or without food.

Withdrawal symptoms seen on discontinuation of sertraline
Abrupt discontinuation should be avoided. When stopping treatment with sertraline the dose should be gradually reduced over a period of at least one to two weeks in order to reduce the risk of withdrawal reactions (see sections 4.4 and 4.8). 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.

4.3 Contraindications

Hypersensitivity to the active substance or any of the excipients listed in section 6.1. Concomitant treatment with irreversible monoamine oxidase inhibitors (MAOIs) is contraindicated due to the risk of serotonin syndrome with symptoms such as agitation, tremor and hyperthermia. Sertraline must not be initiated for at least 14 days after discontinuation of treatment with an irreversible MAOI. Sertraline must be discontinued for at least 7 days before starting treatment with an irreversible MAOI (see section 4.5).

Concomitant intake of pimozide is contraindicated (see section 4.5).

4.4 Special warnings and precautions for use

Serotonin Syndrome (SS) or Neuroleptic Malignant Syndrome (NMS)
The development of potentially life-threatening syndromes like serotonin syndrome (SS) or Neuroleptic Malignant Syndrome (NMS) has been reported with SSRIs, including treatment with sertraline. The risk of SS or NMS with SSRIs is increased with concomitant use of other serotonergic drugs (including other serotonergic antidepressants, amphetamines, triptans), with drugs which impair metabolism of serotonin (including MAOIs e.g. methylene blue), antipsychotics and other dopamine antagonists, and with opiate drugs. Patients should be monitored for the emergence of signs and symptoms of SS or NMS syndrome (see section 4.3).

Switching from Selective Serotonin Reuptake Inhibitors (SSRIs), antidepressants or anti-obsessional drugs
There is limited controlled experience regarding the optimal timing of switching from SSRIs, antidepressants or anti-obsessional drugs to sertraline. Care and prudent medical judgment should be exercised when switching, particularly from long-acting agents such as fluoxetine.

Other serotonergic drugs e.g. tryptophan, fenfluramine and 5-HT agonists
Co-administration of sertraline with other drugs which enhance the effects of serotonergic neurotransmission such as amphetamines, tryptophan or fenfluramine or 5-HT agonists, or the herbal medicine, St John’s Wort (Hypericum perforatum), should be undertaken with caution and avoided whenever possible due to the potential for a pharmacodynamic interaction.
QTc Prolongation/Torsade de Pointes (TdP)
Cases of QTc prolongation and TdP have been reported during post-marketing use of sertraline. The majority of reports occurred in patients with other risk factors for QTc prolongation/TdP. Effect on QTc prolongation was confirmed in a thorough QTc study in healthy volunteers, with a statistically significant positive exposure response relationship. Therefore sertraline should be used with caution in patients with additional risk factors for QTc prolongation such as cardiac disease, hypokalaemia or hypomagnesemia, familial history of QTc prolongation, bradycardia and concomitant use of medications which prolong QTc interval (see sections 4.5 and 5.1).

Activation of hypomania or mania
Manic/hypomanic symptoms have been reported to emerge in a small proportion of patients treated with marketed antidepressant and anti-obsessional drugs, including sertraline. Therefore sertraline should be used with caution in patients with a history of mania/hypomania. Close surveillance by the physician is required. Sertraline should be discontinued in any patient entering a manic phase.

Schizophrenia
Psychotic symptoms might become aggravated in schizophrenic patients.

Seizures
Seizures may occur with sertraline therapy: sertraline should be avoided in patients with unstable epilepsy and patients with controlled epilepsy should be carefully monitored. Sertraline should be discontinued in any patient who develops seizures.

Suicide/suicidal thoughts/suicide attempts or clinical worsening
Depression is associated with an increased risk of suicidal thoughts, self harm and suicide (suicide-related events). This risk persists until significant remission occurs. As improvement may not occur during the first few weeks or more of treatment, patients should be closely monitored until such improvement occurs. It is general clinical experience that the risk of suicide may increase in the early stages of recovery.

Patients with a history of suicide-related events, or those exhibiting a significant degree of suicidal ideation prior to commencement of treatment are known to be at greater risk of suicidal thoughts or suicide attempts, and should receive careful monitoring during treatment. A meta-analysis of placebo-controlled clinical trials of antidepressant drugs in adult patients with psychiatric disorders showed an increased risk of suicidal behaviour with antidepressants compared to placebo in patients less than 25 years old.

Close supervision of patients and in particular those at high risk should accompany drug therapy especially in early treatment and following dose changes. Patients (and caregivers of patients) should be alerted about the need to monitor for any clinical worsening, suicidal behaviour or thoughts and unusual changes in behaviour and to seek medical advice immediately if these symptoms present.

Sexual dysfunction
Selective serotonin reuptake inhibitors (SSRIs) may cause symptoms of sexual dysfunction (see section 4.8). There have been reports of long-lasting sexual dysfunction where the symptoms have continued despite discontinuation of SSRIs.
**Paediatric population**
Sertraline should not be used in the treatment of children and adolescents under the age of 18 years. Suicide-related behaviours (suicide attempt and suicidal thoughts), and hostility (predominantly aggression, oppositional behaviour and anger) were more frequently observed in clinical trials among children and adolescents treated with antidepressants compared to those treated with placebo. If, based on clinical need, a decision to treat is nevertheless taken; the patient should be carefully monitored for appearance of suicidal symptoms. In addition only limited clinical evidence is available concerning, long-term safety data in children and adolescents including effects on growth, sexual maturation and cognitive and behavioural developments. A few cases of retarded growth and delayed puberty have been reported post-marketing. The clinical relevance and causality are yet unclear (see section 5.3 for corresponding preclinical safety data). Physicians must monitor paediatric patients on long term treatment for abnormalities in growth and development.

**Abnormal bleeding/Haemorrhage**
There have been reports of bleeding abnormalities with SSRIs including cutaneous bleeding (ecchymoses and purpura) and other haemorrhagic events such as gastrointestinal or gynaecological bleeding, including fatal haemorrhages. Caution is advised in patients taking SSRIs, particularly in concomitant use with drugs known to affect platelet function (e.g. anticoagulants, atypical antipsychotics and phenothiazines, most tricyclic antidepressants, acetylsalicylic acid and non-steroidal anti-inflammatory drugs (NSAIDs)) as well as in patients with a history of bleeding disorders (see section 4.5).

**Hyponatraemia**
Hyponatraemia may occur as a result of treatment with SSRIs or SNRIs including sertraline. In many cases, hyponatraemia appears to be the result of a syndrome of inappropriate antiuretic hormone secretion (SIADH). Cases of serum sodium levels lower than 110 mmol/L have been reported.
Elderly patients may be at greater risk of developing hyponatraemia with SSRIs and SNRIs. Also patients taking diuretics or who are otherwise volume-depleted may be at greater risk (see Use in elderly). Discontinuation of sertraline should be considered in patients with symptomatic hyponatraemia and appropriate medical intervention should be instituted. Signs and symptoms of hyponatraemia include headache, difficulty concentrating, memory impairment, confusion, weakness and unsteadiness which may lead to falls. Signs and symptoms associated with more severe and/or acute cases have included hallucination, syncope, seizure, coma, respiratory arrest, and death.

**Withdrawal symptoms seen on discontinuation of sertraline treatment**
Withdrawal symptoms when treatment is discontinued are common, particularly if discontinuation is abrupt (see section 4.8). In clinical trials, among patients treated with sertraline, the incidence of reported withdrawal reactions was 23% in those discontinuing sertraline compared to 12% in those who continued to receive sertraline treatment.

The risk of withdrawal symptoms may be dependent on several factors including the duration and dose of therapy and the rate of dose reduction. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances (including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor and headache are the most commonly reported reactions. Generally these symptoms are mild to moderate; however, in some patients they may be severe
in intensity. They usually occur within the first few days of discontinuing treatment, but there have been very rare reports of such symptoms in patients who have inadvertently missed a dose. Generally these symptoms are self-limiting and usually resolve within 2 weeks, though in some individuals they may be prolonged (2-3 months or more). It is therefore advised that sertraline should be gradually tapered when discontinuing treatment over a period of several weeks or months, according to the patient’s needs (see section 4.2).

**Akathisia/psychomotor restlessness**
The use of sertraline has been associated with the development of akathisia, characterised by a subjectively unpleasant or distressing restlessness and need to move often accompanied by an inability to sit or stand still. This is most likely to occur within the first few weeks of treatment. In patients who develop these symptoms, increasing the dose may be detrimental.

**Hepatic impairment**
Sertraline is extensively metabolised by the liver. A multiple dose pharmacokinetic study in subjects with mild, stable cirrhosis demonstrated a prolonged elimination half life and approximately three-fold greater AUC and Cmax in comparison to normal subjects. There were no significant differences in plasma protein binding observed between the two groups. The use of sertraline in patients with hepatic disease must be approached with caution. If sertraline is administered to patients with hepatic impairment, a lower or less frequent dose should be considered. Sertraline should not be used in patients with severe hepatic impairment (see section 4.2).

**Renal impairment**
Sertraline is extensively metabolised, and excretion of unchanged drug in urine is a minor route of elimination. In studies of patients with mild to moderate renal impairment (creatinine clearance 30-60 ml/min) or moderate to severe renal impairment (creatinine clearance 10-29 ml/min), multiple-dose pharmacokinetic parameters (AUC0-24 or Cmax) were not significantly different compared with controls. Sertraline dosing does not have to be adjusted based on the degree of renal impairment.

**Use in elderly**
Over 700 elderly patients (>65 years) have participated in clinical studies. The pattern and incidence of adverse reactions in the elderly was similar to that in younger patients.

SSRIs or SNRIs including sertraline have however been associated with cases of clinically significant hyponatraemia in elderly patients, who may be at greater risk for this adverse event (see Hyponatraemia in section 4.4).

**Diabetes**
In patients with diabetes, treatment with an SSRI may alter glycaemic control. Insulin and/or oral hypoglycaemic dosage may need to be adjusted.

**Electroconvulsive therapy**
There are no clinical studies establishing the risks or benefits of the combined use of ECT and sertraline.

**Grapefruit juice**
The administration of sertraline with grapefruit juice is not recommended (see section 4.5).
Interference with urine screening tests
False-positive urine immunoassay screening tests for benzodiazepines have been reported in patients taking sertraline. This is due to lack of specificity of the screening tests. False-positive test results may be expected for several days following discontinuation of sertraline therapy. Confirmatory tests, such as gas chromatography/mass spectrometry, will distinguish sertraline from benzodiazepines.

Angle-Closure glaucoma
SSRIs including sertraline may have an effect on pupil size resulting in mydriasis. This mydriatic effect has the potential to narrow the eye angle resulting in increased intraocular pressure and angle-closure glaucoma, especially in patients pre-disposed. Sertraline should therefore be used with caution in patients with angle-closure glaucoma or history of glaucoma.

4.5 Interaction with other medicinal products and other forms of interaction

Contraindicated

Monoamine Oxidase Inhibitors
Irreversible MAOIs (e.g. selegiline)
Sertraline must not be used in combination with irreversible MAOIs such as selegiline. Sertraline must not be initiated for at least 14 days after discontinuation of treatment with an irreversible MAOI. Sertraline must be discontinued for at least 7 days before starting treatment with an irreversible MAOI (see section 4.3).

Reversible, selective MAO-A inhibitor (moclobemide)
Due to the risk of serotonin syndrome, the combination of sertraline with a reversible and selective MAOI, such as moclobemide, should not be given. Following treatment with a reversible MAO-inhibitor, a shorter withdrawal period than 14 days may be used before initiation of sertraline treatment. It is recommended that sertraline should be discontinued for at least 14 days before starting treatment with a reversible MAOI (see section 4.3).

Reversible, non-selective MAOI (linezolid)
The antibiotic linezolid is a weak reversible and non-selective MAOI and should not be given to patients treated with sertraline (see section 4.3).

Severe adverse reactions have been reported in patients who have recently been discontinued from an MAOI (e.g. methylene blue) and started on sertraline, or have recently had sertraline therapy discontinued prior to initiation of an MAOI. These reactions have included tremor, myoclonus, diaphoresis, nausea, vomiting, flushing, dizziness, and hyperthermia with features resembling neuroleptic malignant syndrome, seizures, and death.

Pimozide
Increased pimozide levels of approximately 35% have been demonstrated in a study of a single low dose pimozide (2 mg). These increased levels were not associated with any changes in EKG. While the mechanism of this interaction is unknown, due to the narrow therapeutic index of pimozide, concomitant administration of sertraline and pimozide is contraindicated (see section 4.3).
Co-administration with sertraline is not recommended

CNS depressants and alcohol
The co-administration of sertraline 200 mg daily did not potentiate the effects of alcohol, carbamazepine, haloperidol, or phenytoin on cognitive and psychomotor performance in healthy subjects; however, the concomitant use of sertraline and alcohol is not recommended.

Other serotonergic drugs
See section 4.4.

Caution is also advised with fentanyl (used in general anaesthesia or in the treatment of chronic pain), other serotonergic drugs (including other serotonergic antidepressants, amphetamines, triptans), and with other opiate drugs.

Special Precautions

Drugs that Prolong the QT Interval
The risk of QTc prolongation and/or ventricular arrhythmias (e.g., TdP) may be increased with concomitant use of other drugs which prolong the QTc interval (e.g., some antipsychotics and antibiotics) (see sections 4.4 and 5.1).

Lithium
In a placebo-controlled trial in normal volunteers, the co-administration of sertraline with lithium did not significantly alter lithium pharmacokinetics, but did result in an increase in tremor relative to placebo, indicating a possible pharmacodynamic interaction. When co-administering sertraline with lithium, patients should be appropriately monitored.

Phenytoin
A placebo-controlled trial in normal volunteers suggests that chronic administration of sertraline 200 mg/day does not produce clinically important inhibition of phenytoin metabolism. Nonetheless, as some case reports have emerged of high phenytoin exposure in patients using sertraline, it is recommended that plasma phenytoin concentrations be monitored following initiation of sertraline therapy, with appropriate adjustments to the phenytoin dose. In addition, co-administration of phenytoin may cause a reduction of sertraline plasma levels. It cannot be excluded that other CYP3A4 inducers, e.g. phenobarbital, carbamazepine, St John’s Wort, rifampicin may cause a reduction of sertraline plasma levels.

Triptans
There have been rare post-marketing reports describing patients with weakness, hyperreflexia, incoordination, confusion, anxiety and agitation following the use of sertraline and sumatriptan. Symptoms of serotonergic syndrome may also occur with other products of the same class (triptans). If concomitant treatment with sertraline and triptans is clinically warranted, appropriate observation of the patient is advised (see section 4.4).

Warfarin
Co-administration of sertraline 200 mg daily with warfarin resulted in a small but statistically significant increase in prothrombin time, which may in some rare cases unbalance the INR value.
Accordingly, prothrombin time should be carefully monitored when sertraline therapy is initiated or stopped.

**Other drug interactions, digoxin, atenolol, cimetidine**

Co-administration with cimetidine caused a substantial decrease in sertraline clearance. The clinical significance of these changes is unknown. Sertraline had no effect on the beta-adrenergic blocking ability of atenolol. No interaction of sertraline 200 mg daily was observed with digoxin.

**Drugs affecting platelet function**

The risk of bleeding may be increased when medicines acting on platelet function (e.g. NSAIDs, acetylsalicylic acid and ticlopidine) or other medicines that might increase bleeding risk are concomitantly administered with SSRIs, including sertraline (see section 4.4).

**Neuromuscular Blockers**

SSRIs may reduce plasma cholinesterase activity resulting in a prolongation of the neuromuscular blocking action of mivacurium or other neuromuscular blockers.

**Drugs Metabolized by Cytochrome P450**

Sertraline may act as a mild-moderate inhibitor of CYP 2D6. Chronic dosing with sertraline 50 mg daily showed moderate elevation (mean 23%-37%) of steady-state desipramine plasma levels (a marker of CYP 2D6 isozyme activity). Clinical relevant interactions may occur with other CYP 2D6 substrates with a narrow therapeutic index like class 1C antiarrhythmics such as propafenone and flecainide, TCAs and typical antipsychotics, especially at higher sertraline dose levels.

Sertraline does not act as an inhibitor of CYP 3A4, CYP 2C9, CYP 2C19, and CYP 1A2 to a clinically significant degree. This has been confirmed by in-vivo interaction studies with CYP3A4 substrates (endogenous cortisol, carbamazepine, terfenadine, alprazolam), CYP2C19 substrate diazepam, and CYP2C9 substrates tolbutamide, glibenclamide and phenytoin. In vitro studies indicate that sertraline has little or no potential to inhibit CYP 1A2.

Intake of three glasses of grapefruit juice daily increased the sertraline plasma levels by approximately 100% in a cross-over study in eight Japanese healthy subjects. Therefore, the intake of grapefruit juice should be avoided during treatment with sertraline (see section 4.4).

Based on the interaction study with grapefruit juice, it cannot be excluded that the concomitant administration of sertraline and potent CYP3A4 inhibitors, e.g. protease inhibitors, ketoconazole, itraconazole, posaconazole, voriconazole, clarithromycin, telithromycin and nefazodone, would result in even larger increases in exposure of sertraline. This also concerns moderate CYP3A4 inhibitors, e.g. aprepitant, erythromycin, fluconazole, verapamil and diltiazem. The intake of potent CYP3A4 inhibitors should be avoided during treatment with sertraline.

Sertraline plasma levels are enhanced by about 50% in poor metabolizers of CYP2C19 compared to rapid metabolizers (see section 5.2). Interaction with strong inhibitors of CYP2C19, e.g. omeprazole, lansoprazole, pantoprazole, rabeprazole, fluoxetine, fluvoxamine cannot be excluded.
4.6  Fertility, pregnancy and lactation

Pregnancy

There are no well controlled studies in pregnant women. However, a substantial amount of data did not reveal evidence of induction of congenital malformations by sertraline. Animal studies showed evidence for effects on reproduction probably due to maternal toxicity caused by the pharmacodynamic action of the compound and/or direct pharmacodynamic action of the compound on the foetus (see section 5.3).

Use of sertraline during pregnancy has been reported to cause symptoms, compatible with withdrawal reactions, in some neonates, whose mothers had been on sertraline. This phenomenon has also been observed with other SSRI antidepressants. Sertraline is not recommended in pregnancy, unless the clinical condition of the woman is such that the benefit of the treatment is expected to outweigh the potential risk.

Neonates should be observed if maternal use of sertraline continues into the later stages of pregnancy, particularly the third trimester. The following symptoms may occur in the neonate after maternal sertraline use in later stages of pregnancy: respiratory distress, cyanosis, apnoea, seizures, temperature instability, feeding difficulty, vomiting, hypoglycaemia, hypertonia, hypotonia, hyperreflexia, tremor, jitteriness, irritability, lethargy, constant crying, somnolence and difficulty in sleeping. These symptoms could be due to either serotonergic effects or withdrawal symptoms. In a majority of instances the complications begin immediately or soon (<24 hours) after delivery.

Epidemiological data have suggested that the use of SSRIs in pregnancy, particular in late pregnancy, may increase the risk of persistent pulmonary hypertension in the newborn (PPHN). The observed risk was approximately 5 cases per 1000 pregnancies. In the general population 1 to 2 cases of PPHN per 1000 pregnancies occur.

Breast-feeding

Published data concerning sertraline levels in breast milk show that small quantities of sertraline and its metabolite N-desmethylsertraline are excreted in milk. Generally negligible to undetectable levels were found in infant serum, with one exception of an infant with serum levels about 50% of the maternal level (but without a noticeable health effect in this infant). To date, no adverse effects on the health of infants nursed by mothers using sertraline have been reported, but a risk cannot be excluded. Use in nursing mothers is not recommended unless, in the judgment of the physician, the benefit outweighs the risk.

Fertility

Animal data did not show an effect of sertraline on fertility parameters (see section 5.3.). Human case reports with some SSRI's have shown that an effect on sperm quality is reversible. Impact on human fertility has not been observed so far.

4.7  Effects on ability to drive and use machines

Clinical pharmacology studies have shown that sertraline has no effect on psychomotor performance. However, as psychotrope drugs may impair the mental or physical abilities required for the performance of potentially hazardous tasks such as driving a car or operating machinery, the patient should be cautioned accordingly.
4.8 Undesirable effects

Nausea is the most common undesirable effect. In the treatment of social anxiety disorder, sexual dysfunction (ejaculation failure) in men occurred in 14% for sertraline vs 0% in placebo. These undesirable effects are dose dependent and are often transient in nature with continued treatment.

Table 1 displays adverse reactions observed from postmarketing experience (frequency not known) and placebo-controlled clinical trials (comprising a total of 2542 patients on sertraline and 2145 on placebo) in depression, OCD, panic disorder, PTSD and social anxiety disorder. Some adverse drug reactions listed in Table 1 may decrease in intensity and frequency with continued treatment and do not generally lead to cessation of therapy.

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Very Common (≥1/10)</th>
<th>Common (≥1/100 to &lt;1/10)</th>
<th>Uncommon (≥1/1,000 to &lt;1/100)</th>
<th>Rare (≥1/10,000 to &lt;1/1,000)</th>
<th>Frequency Not Known (Cannot be Estimated From the Available Data)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infections and infestations</td>
<td>upper respiratory tract infection, pharyngitis, rhinitis</td>
<td>gastroenteritis, otitis media</td>
<td></td>
<td>diverticulitis§</td>
<td></td>
</tr>
<tr>
<td>Neoplasms benign, malignant and unspecified (including cysts and polyps)</td>
<td></td>
<td></td>
<td></td>
<td>neoplasm</td>
<td></td>
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<td>Blood and lymphatic system disorders</td>
<td></td>
<td></td>
<td></td>
<td>lymphadenopathy, thrombocytopenia*§, leukopenia*§</td>
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<td>Immune system disorders</td>
<td>hypersensitivity*, seasonal allergy*</td>
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<td></td>
<td>anaphylactoid reaction*</td>
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<td>Endocrine disorders</td>
<td></td>
<td></td>
<td></td>
<td>hyperprolactinaemia* §, inappropriate antidiuretic hormone secretion*§</td>
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</tr>
<tr>
<td>Metabolism and nutrition disorders</td>
<td>decreased appetite, increased appetite*</td>
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<td></td>
<td>hypercholesterolaemia, diabetes mellitus*, hypoglycaemia*, hyperglycaemia*§, hyponatraemia*§</td>
<td></td>
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<tr>
<td>Psychiatric</td>
<td>insomnia anxiety*, suicidal</td>
<td></td>
<td></td>
<td>conversion</td>
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</tbody>
</table>
### Table 1: Adverse Reactions

Frequency of adverse reactions observed from placebo-controlled clinical trials in depression, OCD, panic disorder, PTSD and social anxiety disorder. Pooled analysis and post-marketing experience.

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<th>System Organ Class</th>
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<tr>
<td>disorders</td>
<td>depression*, agitation*, libido decreased*, nervousness, depersonalisation, nightmare, bruxism*</td>
<td>ideation/behaviour, psychotic disorder*, thinking abnormal, apathy, hallucination*, aggression*, euphoric mood*, paranoia</td>
<td>disorder*, paroniria*, drug dependence, sleep walking, premature ejaculation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>dizziness, headache*, somnolence</td>
<td>tremor, movement disorders (including extrapyramidal symptoms such as hyperkinesia, hypertonia, dystonia, teeth grinding or gait abnormalities), paraesthesia*, hypertonia*, disturbance in attention, dysgeusia</td>
<td>amnesia, hypoaesthesia*, muscle contractions involuntary*, syncope*, hyperkinesia*, migraine*, convulsion*, dizziness postural, coordination abnormal, speech disorder</td>
<td>coma*, akathisia (see section 4.4), dyskinesia, hyperaesthesia, cerebrovascular spasm (including reversible cerebral vasoconstriction syndrome and Call-Fleming syndrome)<em>, psychomotor restlessness</em> (see section 4.4), sensory disturbance, choreoathetosis*, also reported were signs and symptoms associated with serotonin syndrome* or neuroleptic malignant syndrome: In some cases associated with concomitant use of serotonergic drugs that included agitation, confusion,</td>
<td></td>
</tr>
</tbody>
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</tr>
</thead>
<tbody>
<tr>
<td>Eye disorders</td>
<td>Diaphoresis, diarrhoea, fever, hypertension, rigidity and tachycardia&lt;sup&gt;8&lt;/sup&gt;</td>
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<td>Ear and labyrinth disorders</td>
<td>Tinnitus&lt;sup&gt;*&lt;/sup&gt;</td>
<td>Ear pain</td>
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<td></td>
<td></td>
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<tr>
<td>Cardiac disorders</td>
<td>Palpitations&lt;sup&gt;*&lt;/sup&gt;</td>
<td>Tachycardia&lt;sup&gt;*&lt;/sup&gt;, cardiac disorder</td>
<td></td>
<td>Myocardial infarction&lt;sup&gt;9&lt;/sup&gt;, Torsade de Pointes&lt;sup&gt;13&lt;/sup&gt; (see sections 4.4, 4.5 and 5.1), bradycardia, QTc prolongation&lt;sup&gt;*&lt;/sup&gt; (see sections 4.4, 4.5 and 5.1)</td>
<td>Maculopathy</td>
</tr>
<tr>
<td>Vascular disorders</td>
<td>Hot flush&lt;sup&gt;*&lt;/sup&gt;</td>
<td>Abnormal bleeding (such as gastrointestinal bleeding)&lt;sup&gt;<em>&lt;/sup&gt;, hypertension&lt;sup&gt;</em>&lt;/sup&gt;, flushing, haematuria&lt;sup&gt;*&lt;/sup&gt;</td>
<td></td>
<td>Peripheral ischaemia</td>
<td></td>
</tr>
<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
<td>Yawning&lt;sup&gt;*&lt;/sup&gt;</td>
<td>Dyspnoea, epistaxis&lt;sup&gt;<em>&lt;/sup&gt;, bronchospasm&lt;sup&gt;</em>&lt;/sup&gt;</td>
<td>Hyperventilation, interstitial lung disease&lt;sup&gt;4&lt;/sup&gt;, laryngospasm, dysphonia, stridor&lt;sup&gt;*&lt;/sup&gt;, hypoventilation, hiccups</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gastrointestinal disorders</td>
<td>Nausea, diarrhoea, constipation&lt;sup&gt;<em>&lt;/sup&gt;, abdominal pain&lt;sup&gt;</em>&lt;/sup&gt;, vomiting&lt;sup&gt;*&lt;/sup&gt;, flatulence</td>
<td>Melaena, tooth disorder, oesophagitis, glossitis, haemorrhoids, mouth ulceration, pancreatitis&lt;sup&gt;4&lt;/sup&gt;, haematochezia, tongue ulceration, stomatitis</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 1: Adverse Reactions
Frequency of adverse reactions observed from placebo-controlled clinical trials in depression, OCD, panic disorder, PTSD and social anxiety disorder. Pooled analysis and post-marketing experience.

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Very Common (≥1/10)</th>
<th>Common (≥1/100 to &lt;1/10)</th>
<th>Uncommon (≥1/1,000 to &lt;1/100)</th>
<th>Rare (≥1/10,000 to &lt;1/1,000)</th>
<th>Frequency Not Known (Cannot be Estimated From the Available Data)</th>
</tr>
</thead>
<tbody>
<tr>
<td>salivary hypersecretion, dysphagia, eructation, tongue disorder</td>
<td>hepatic function abnormal, serious liver events (including hepatitis, jaundice and hepatic failure)</td>
<td>rare reports of severe cutaneous adverse reactions (SCAR): e.g. Stevens-Johnson syndrome*, epidermal necrolysis§, skin reaction§, photosensitivity§, angioedema, hair texture abnormal, skin odour abnormal, dermatitis bullous, rash follicular</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Skin and subcutaneous tissue disorders</td>
<td>hyperhidrosis, rash*</td>
<td>periorbital oedema*, urticaria*, alopecia*, pruritus*, purpura*, dermatitis, dry skin, face oedema, cold sweat</td>
<td>rare reports of severe cutaneous adverse reactions (SCAR): e.g. Stevens-Johnson syndrome*, epidermal necrolysis§, skin reaction§, photosensitivity§, angioedema, hair texture abnormal, skin odour abnormal, dermatitis bullous, rash follicular</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>back pain, arthralgia*, myalgia</td>
<td>osteoarthritis, muscle twitching, muscle cramps*, muscular weakness</td>
<td>rhabdomyolysis§, bone disorder</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Renal and urinary disorders</td>
<td>pollakiuria, micturition disorder, urinary retention, urinary incontinence*</td>
<td></td>
<td>urinary hesitation*, oliguria</td>
<td></td>
<td></td>
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<tr>
<td>Reproductive system and breast disorders</td>
<td>ejaculatio n failure</td>
<td>menstruation irregular*, erectile dysfunction</td>
<td>sexual dysfunction (see section 4.4), menorrhagia, vaginal haemorrhage, female sexual dysfunction-(see section 4.4)</td>
<td>galactorrhoea*, atrophic vulvovaginitis, genital discharge, balanoposthitis*, gynaecomastia*, priapism*</td>
<td></td>
</tr>
<tr>
<td>General disorders and administration site conditions</td>
<td>fatigue*</td>
<td>malaise*, chest pain*, asthenia*, pyrexia*</td>
<td>oedema peripheral*, chills, gait disturbance*, thirst</td>
<td>hernia, drug tolerance decreased</td>
<td></td>
</tr>
<tr>
<td>Investigations</td>
<td>weight increased*</td>
<td>alanine aminotransferase increased*, aspartate aminotransferase increased*, weight decreased*</td>
<td>blood cholesterol increased*, abnormal clinical laboratory results, semen abnormal, altered platelet function*§</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Injury, poisoning and procedural complications</td>
<td>injury</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surgical and medical procedures</td>
<td></td>
<td></td>
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</tbody>
</table>

* ADR identified post-marketing
§ ADR frequency represented by the estimated upper limit of the 95% confidence interval using “The Rule of 3”.

Withdrawal symptoms seen on discontinuation of sertraline treatment
Discontinuation of sertraline (particularly when abrupt) commonly leads to withdrawal symptoms. Dizziness, sensory disturbances (including paraesthesia), sleep disturbances
(including insomnia and intense dreams), agitation or anxiety, nausea and/or vomiting, tremor and headache are the most commonly reported. Generally these events are mild to moderate and are self-limiting; however, in some patients they may be severe and/or prolonged. It is therefore advised that when sertraline treatment is no longer required, gradual discontinuation by dose tapering should be carried out (see sections 4.2 and 4.4).

**Elderly population**

SSRIs or SNRIs including sertraline have been associated with cases of clinically significant hyponatraemia in elderly patients, who may be at greater risk for this adverse event (see section 4.4).

**Paediatric population**

In over 600 paediatric patients treated with sertraline, the overall profile of adverse reactions was generally similar to that seen in adult studies. The following adverse reactions were reported from controlled trials (n=281 patients treated with sertraline):

- **Very common (≥1/10)**: Headache (22%), insomnia (21%), diarrhoea (11%) and nausea (15%).
- **Common (≥1/100 to <1/10)**: Chest pain, mania, pyrexia, vomiting, anorexia, affect lability, aggression, agitation, nervousness, disturbance in attention, dizziness, hyperkinesia, migraine, somnolence, tremor, visual disturbance, dry mouth, dyspepsia, nightmare, fatigue, urinary incontinence, rash, acne, epistaxis, flatulence.
- **Uncommon (≥1/1000 to <1/100)**: ECG QT prolonged (see sections 4.4, 4.5 and 5.1), suicide attempt, convulsion, extrapyramidal disorder, paraesthesia, depression, hallucination, purpura, hyperventilation, anaemia, hepatic function abnormal, alanine aminotransferase increased, cystitis, herpes simplex, otitis externa, ear pain, eye pain, mydriasis, malaise, haematuria, rash pustular, rhinitis, injury, weight decreased, muscle twitching, abnormal dreams, apathy, albuminuria, pollakiuria, polyuria, breast pain, menstrual disorder, alopecia, dermatitis, skin disorder, skin odour abnormal, urticaria, bruxism, flushing.

*Frequency not known*: enuresis

**Class effects**

Epidemiological studies, mainly conducted in patients 50 years of age and older, show an increased risk of bone fractures in patients receiving SSRIs and TCAs. The mechanism leading to this risk is unknown.

**Reporting of suspected adverse reactions**

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product.

Any suspected adverse events should be reported to the Ministry of Health according to the National Regulation by using an online form https://sideeffects.health.gov.il/

**4.9 Overdose**

**Toxicity**

Sertraline has a margin of safety dependent on patient population and/or concomitant medication. Deaths have been reported involving overdoses of sertraline, alone or in combination with other drugs and/or alcohol. Therefore, any overdosage should be medically treated aggressively.
Symptoms
Symptoms of overdose include serotonin-mediated side effects such as somnolence, gastrointestinal disturbances (e.g. nausea and vomiting), tachycardia, tremor, agitation and dizziness. Coma has been reported although less frequently.

QTc prolongation/Torsade de Pointes has been reported following sertraline overdose; therefore, ECG-monitoring is recommended in all ingestions of sertraline overdoses (see sections 4.4, 4.5 and 5.1).

Management
There are no specific antidotes to sertraline. It is recommended to establish and maintain an airway and, if necessary, ensure adequate oxygenation and ventilation. Activated charcoal, which may be used with a cathartic, may be as or more effective than lavage, and should be considered in treating overdose. Induction of emesis is not recommended. Cardiac (e.g. ECG) and vital sign monitoring is also recommended, along with general symptomatic and supportive measures. Due to the large volume of distribution of sertraline, forced diuresis, dialysis, haemoperfusion and exchange transfusion are unlikely to be of benefit.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Selective serotonin reuptake inhibitors (SSRI), ATC code: N06 AB06.

Mechanism of action

Sertraline is a potent and specific inhibitor of neuronal serotonin (5 HT) uptake in vitro, which results in the potentiation of the effects of 5-HT in animals. It has only very weak effects on norepinephrine and dopamine neuronal reuptake. At clinical doses, sertraline blocks the uptake of serotonin into human platelets. It is devoid of stimulant, sedative or anticholinergic activity or cardiotoxicity in animals. In controlled studies in normal volunteers, sertraline did not cause sedation and did not interfere with psychomotor performance. In accord with its selective inhibition of 5-HT uptake, sertraline does not enhance catecholaminergic activity. Sertraline has no affinity for muscarinic (cholinergic), serotonergic, dopaminergic, adrenergic, histaminergic, GABA or benzodiazepine receptors. The chronic administration of sertraline in animals was associated with down-regulation of brain norepinephrine receptors as observed with other clinically effective antidepressants and antiobsessional drugs.

Sertraline has not demonstrated potential for abuse. In a placebo-controlled, double-blind randomized study of the comparative abuse liability of sertraline, alprazolam and d-amphetamine in humans, sertraline did not produce positive subjective effects indicative of abuse potential. In contrast, subjects rated both alprazolam and d-amphetamine significantly greater than placebo on measures of drug liking, euphoria and abuse potential. Sertraline did not produce either the stimulation and anxiety associated with d-amphetamine or the sedation and psychomotor impairment associated with alprazolam. Sertraline does not function as a
positive reinforcer in rhesus monkeys trained to self administer cocaine, nor does it substitute as a discriminative stimulus for either d-amphetamine or pentobarbital in rhesus monkeys.

**Clinical efficacy and safety**

**Major Depressive Disorder**

A study was conducted which involved depressed outpatients who had responded by the end of an initial 8-week open treatment phase on sertraline 50-200 mg/day. These patients (n=295) were randomized to continuation for 44 weeks on double-blind sertraline 50-200 mg/day or placebo. A statistically significantly lower relapse rate was observed for patients taking sertraline compared to those on placebo. The mean dose for completers was 70 mg/day. The % of responders (defined as those patients that did not relapse) for sertraline and placebo arms were 83.4% and 60.8%, respectively.

**Cardiac Electrophysiology**

In a dedicated thorough QTc study, conducted at steady state at supratherapeutic exposures in healthy volunteers (treated with 400 mg/day, twice the maximum recommended daily dose), the upper bound of the 2-sided 90% CI for the time matched Least Square mean difference of QTcF between sertraline and placebo (11.666 msec) was greater than the predefined threshold of 10 msec at the 4-hour postdose time point. Exposure-response analysis indicated a slightly positive relationship between QTcF and sertraline plasma concentrations [0.036 msec/(ng/mL); p<0.0001]. Based on the exposure response model, the threshold for clinically significant prolongation of the QTcF (i.e. for predicted 90% CI to exceed 10 msec) is at least 2.6-fold greater than the average Cmax (86 ng/mL) following the highest recommended dose of sertraline (200 mg/day) (see sections 4.4, 4.5, 4.8 and 4.9).

**5.2 Pharmacokinetic properties**

**Absorption**

In man, following an oral once-daily dosage of 50 to 200 mg for 14 days, peak plasma concentrations of sertraline occur at 4.5 to 8.4 hours after the daily administration of the drug. Food does not significantly change the bioavailability of sertraline tablets.

**Distribution**

Approximately 98% of the circulating drug is bound to plasma proteins.

**Biotransformation**

Sertraline undergoes extensive first-pass hepatic metabolism.

Based on clinical and *in-vitro* data, it can be concluded that sertraline is metabolized by multiple pathways including CYP3A4, CYP2C19 (see section 4.5) and CYP2B6. Sertraline and its major metabolite desmethylsertraline are also substrate of P-glycoprotein *in-vitro*.

**Elimination**

The mean half-life of sertraline is approximately 26 hours (range 22-36 hours). Consistent with the terminal elimination half-life, there is an approximately two-fold accumulation up to steady state concentrations, which are achieved after one week of once-daily dosing. The half-life of N-desmethylsertraline is in the range of 62 to 104 hours. Sertraline and N-desmethylsertraline are both extensively metabolized in man and the resultant metabolites excreted in faeces and urine in equal amounts. Only a small amount (<0.2%) of unchanged sertraline is excreted in the urine.
**Linearity/non-linearity**
Sertraline exhibits dose proportional pharmacokinetics in the range of 50 to 200 mg.

**Pharmacokinetics in specific patient groups**

**Adolescents and elderly**
The pharmacokinetic profile in adolescents or elderly is not significantly different from that in adults between 18 and 65 years.

**Hepatic impairment**
In patients with liver damage, the half life of sertraline is prolonged and AUC is increased three fold (see sections 4.2 and 4.4).

**Renal impairment**
In patients with moderate-severe renal impairment, there was no significant accumulation of sertraline.

**Pharmacogenomics**
Plasma levels of sertraline were about 50% higher in poor metabolizers of CYP2C19 versus extensive metabolizers. The clinical meaning is not clear, and patients need to be titrated based on clinical response.

### 5.3 Preclinical safety data

Preclinical data does not indicate any special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity and carcinogenesis. Reproduction toxicity studies in animals showed no evidence of teratogenicity or adverse effects on male fertility. Observed foetotoxicity was probably related to maternal toxicity. Postnatal pup survival and body weight were decreased only during the first days after birth. Evidence was found that the early postnatal mortality was due to in-utero exposure after day 15 of pregnancy. Postnatal developmental delays found in pups from treated dams were probably due to effects on the dams and therefore not relevant for human risk.

Animal data from rodents and non-rodents does not reveal effects on fertility.

**Juvenile animal studies**

A juvenile toxicity study in rats has been conducted in which sertraline was administered orally to male and female rats on Postnatal Days 21 through 56 (at doses of 10, 40, or 80 mg/kg/day) with a nondosing recovery phase up to Postnatal Day 196. Delays in sexual maturation occurred in males and females at different dose levels (males at 80 mg/kg and females at ≥10 mg/kg), but despite this finding there were no sertraline-related effects on any of the male or female reproductive endpoints that were assessed. In addition, on Postnatal Days 21 to 56, dehydration, chromorrhinorrhea, and reduced average body weight gain was also observed. All of the aforementioned effects attributed to the administration of sertraline were reversed at some point during the nondosing recovery phase of the study. The clinical relevance of these effects observed in rats administered sertraline has not been established.
6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Sertraline film-coated tablets:

*Tablet cores:*  
- microcrystalline cellulose  
- calcium hydrogen phosphate dihydrate  
- sodium starch glycollate  
- hydroxypropyl cellulose  
- magnesium stearate  
- purified water

*Film coating:*  
- White Opadry  
- Clear Opadry

6.2 Incompatibilities

Not applicable.

6.3 Shelf-life

The expiry date of the product is indicated on the packaging materials.

6.4 Special precautions for storage

Store below 25°C.

6.5 Nature and contents of container

Tablets are packed in Aluminium/PVC blisters of 10 or 14 tablets (packages of 20 or 28 tablets).

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

No special requirements.

Any unused product or waste material should be disposed of in accordance with local requirements.

7. MANUFACTURER

Haupt Pharma Latina S.r.l, Latina, Italy  
Or  
Pfizer Manufacturing Deutschland GmbH, Freiburg, Germany.
8. LICENSE HOLDER

Pfizer PFE Pharmaceuticals Israel Ltd., 9 Shenkar St., Herzliya Pituach 46725.

9. LICENSE NUMBER

Lustral® 50mg: 120-51-27480
Lustral® 100mg: 120-52-27481

The content of this leaflet was approved by the Ministry of Health in March 2017 and updated according to the guidelines of the Ministry of Health in January 2020