#### NORTRIPTYLINE HYDROCHLORIDE- nortriptyline hydrochloride capsule Aidarex Pharmaceuticals LLC

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NORTRIPTYLINE HYDROCHLORIDE CAPSULES USP 0810 0811 0812 0813 Rx only

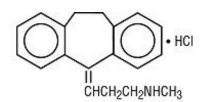
Suicidality and Antidepressant Drugs

Antidepressants increased the risk compared to placebo of suicidal thinking and behavior (suicidality) in children, adolescents, and young adults in short-term studies of major depressive disorder (MDD) and other psychiatric disorders. Anyone considering the use of nortriptyline hydrochloride capsules or any other antidepressant in a child, adolescent, or young adult must balance this risk with the clinical need. Short-term studies did not show an increase in the risk of suicidality with antidepressants compared to placebo in adults beyond age 24; there was a reduction in risk with antidepressants compared to placebo in adults aged 65 and older. Depression and certain other psychiatric disorders are themselves associated with increases in the risk of suicidality. Patients of all ages who are started on antidepressant therapy should be monitored appropriately and observed closely for clinical worsening, suicidality, or unusual changes in behavior. Families and caregivers should be advised of the need for close observation and communication with the prescriber. Nortriptyline hydrochloride capsules are not approved for use in pediatric patients (see WARNINGS, Clinical Worsening and Suicide Risk; PRECAUTIONS, Information for Patients; and PRECAUTIONS, Pediatric Use).

#### DESCRIPTION

Nortriptyline hydrochloride, USP is 1-Propanamine, 3-(10,11-dihydro-5H-dibenzo[a,d]cyclohepten-5-ylidene)-N-methyl-, hydrochloride.

The structural formula is as follows:



C<sub>19</sub>H<sub>21</sub>N•HCl M.W. 299.8

Nortriptyline hydrochloride, USP is a white to off-white powder, having a slight, characteristic odor. It is soluble in water and in chloroform; sparingly soluble in methanol; and practically insoluble in most organic solvents.

Each capsule, for oral administration, contains nortriptyline hydrochloride, USP equivalent to 10 mg, 25 mg, 50 mg or 75 mg nortriptyline.

In addition, each capsule contains the following inactive ingredients: corn starch, D&C Yellow #10 aluminum lake, FD&C Blue #1 aluminum lake, FD&C Blue #2 aluminum lake, FD&C Red #40 aluminum

lake, gelatin, iron oxide black, methylparaben, propylene glycol, propylparaben, shellac glaze, silicone fluid, sodium lauryl sulfate, and titanium dioxide. The 10 mg, 25 mg and 75 mg capsules also contain D&C Yellow #10, FD&C Blue #1, and FD&C Yellow #6.

## CLINICAL PHARMACOLOGY

The mechanism of mood elevation by tricyclic antidepressants is at present unknown. Nortriptyline hydrochloride is not a monoamine oxidase inhibitor. It inhibits the activity of such diverse agents as histamine, 5-hydroxytryptamine, and acetylcholine. It increases the pressor effect of norepinephrine but blocks the pressor response of phenethylamine. Studies suggest that nortriptyline hydrochloride interferes with the transport, release, and storage of catecholamines. Operant conditioning techniques in rats and pigeons suggest that nortriptyline hydrochloride has a combination of stimulant and depressant properties.

#### INDICATIONS AND USAGE

Nortriptyline hydrochloride capsules are indicated for the relief of symptoms of depression. Endogenous depressions are more likely to be alleviated than are other depressive states.

#### CONTRAINDICATIONS

#### Monoamine Oxidase Inhibitors (MAOIs)

The use of MAOIs intended to treat psychiatric disorders with nortriptyline hydrochloride or within 14 days of stopping treatment with nortriptyline hydrochloride is contraindicated because of an increased risk of serotonin syndrome. The use of nortriptyline hydrochloride within 14 days of stopping an MAOI intended to treat psychiatric disorders is also contraindicated (see **WARNINGS** and **DOSAGE AND ADMINISTRATION**).

Starting nortriptyline hydrochloride in a patient who is being treated with MAOIs such as linezolid or intravenous methylene blue is also contraindicated because of an increased risk of serotonin syndrome (see **WARNINGS** and **DOSAGE AND ADMINISTRATION**).

#### Hypersensitivity to Tricyclic Antidepressants

Cross-sensitivity between nortriptyline hydrochloride and other dibenzazepines is a possibility.

## **Myocardial Infarction**

Nortriptyline hydrochloride is contraindicated during the acute recovery period after myocardial infarction.

## WARNINGS

## Clinical Worsening and Suicide Risk

Patients with major depressive disorder (MDD), both adult and pediatric, may experience worsening of their depression and/or the emergence of suicidal ideation and behavior (suicidality) or unusual changes in behavior, whether or not they are taking antidepressant medications, and this risk may persist until significant remission occurs. Suicide is a known risk of depression and certain other psychiatric disorders, and these disorders themselves are the strongest predictors of suicide. There has been a long-standing concern, however, that antidepressants may have a role in inducing worsening of depression and the emergence of suicidality in certain patients during the early phases of treatment. Pooled analyses of short-term placebo-controlled trials of antidepressant drugs (SSRIs and others) showed that these drugs increase the risk of suicidal thinking and behavior (suicidality) in children,

adolescents, and young adults (ages 18 to 24) with major depressive disorder (MDD) and other psychiatric disorders. Short-term studies did not show an increase in the risk of suicidality with antidepressants compared to placebo in adults beyond age 24; there was a reduction with antidepressants compared to placebo in adults.

The pooled analyses of placebo-controlled trials in children and adolescents with MDD, obsessive compulsive disorder (OCD), or other psychiatric disorders included a total of 24 short-term trials of 9 antidepressant drugs in over 4400 patients. The pooled analyses of placebo-controlled trials in adults with MDD or other psychiatric disorders included a total of 295 short-term trials (median duration of 2 months) of 11 antidepressant drugs in over 77,000 patients. There was considerable variation in risk of suicidality among drugs, but a tendency toward an increase in the younger patients for almost all drugs studied. There were differences in absolute risk of suicidality across the different indications, with the highest incidence in MDD. The risk differences (drug vs placebo), however, were relatively stable within age strata and across indications. These risk differences (drug-placebo difference in the number of cases of suicidality per 1000 patients treated) are provided in **Table 1**.

Age Range	Drug-Placebo Difference in Number of Cases of Suicidality per 1000 Patients Treated						
Increases Compared to Placebo							
< 18	14 additional cases						
18 to 24	5 additional cases						
D	ecreases Compared to Placebo						
25 to 64	1 fewer case						
≥ 65	6 fewer cases						

Table 1

No suicides occurred in any of the pediatric trials. There were suicides in the adult trials, but the number was not sufficient to reach any conclusion about drug effect on suicide.

It is unknown whether the suicidality risk extends to longer-term use, i.e., beyond several months. However, there is substantial evidence from placebo-controlled maintenance trials in adults with depression that the use of antidepressants can delay the recurrence of depression.

#### All patients being treated with antidepressants for any indication should be monitored appropriately and observed closely for clinical worsening, suicidality, and unusual changes in behavior, especially during the initial few months of a course of drug therapy, or at times of dose changes, either increases or decreases.

The following symptoms, anxiety, agitation, panic attacks, insomnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomania, and mania, have been reported in adult and pediatric patients being treated with antidepressants for major depressive disorder as well as for other indications, both psychiatric and nonpsychiatric. Although a causal link between the emergence of such symptoms and either the worsening of depression and/or the emergence of suicidal impulses has not been established, there is concern that such symptoms may represent precursors to emerging suicidality.

Consideration should be given to changing the therapeutic regimen, including possibly discontinuing the medication, in patients whose depression is persistently worse, or who are experiencing emergent suicidality or symptoms that might be precursors to worsening depression or suicidality, especially if these symptoms are severe, abrupt in onset, or were not part of the patient's presenting symptoms.

Families and caregivers of patients being treated with antidepressants for major depressive disorder or other indications, both psychiatric and nonpsychiatric, should be alerted about the need to monitor patients for the emergence of agitation, irritability, unusual changes in behavior,

and the other symptoms described above, as well as the emergence of suicidality, and to report such symptoms immediately to health care providers.Such monitoring should include daily observation by families and caregivers. Prescriptions for nortriptyline hydrochloride capsules should be written for the smallest quantity of capsules consistent with good patient management, in order to reduce the risk of overdose.

#### Screening Patients for Bipolar Disorder

A major depressive episode may be the initial presentation of bipolar disorder. It is generally believed (though not established in controlled trials) that treating such an episode with an antidepressant alone may increase the likelihood of precipitation of a mixed/manic episode in patients at risk for bipolar disorder. Whether any of the symptoms described above represent such a conversion is unknown. However, prior to initiating treatment with an antidepressant, patients with depressive symptoms should be adequately screened to determine if they are at risk for bipolar disorder; such screening should include a detailed psychiatric history, including a family history of suicide, bipolar disorder, and depression. It should be noted that nortriptyline hydrochloride capsules are not approved for use in treating bipolar depression.

Patients with cardiovascular disease should be given nortriptyline hydrochloride only under close supervision because of the tendency of the drug to produce sinus tachycardia and to prolong the conduction time. Myocardial infarction, arrhythmia, and strokes have occurred. The antihypertensive action of guanethidine and similar agents may be blocked. Because of its anticholinergic activity, nortriptyline hydrochloride should be used with great caution in patients who have glaucoma or a history of urinary retention. Patients with a history of seizures should be followed closely when nortriptyline hydrochloride is administered, inasmuch as this drug is known to lower the convulsive threshold. Great care is required if nortriptyline hydrochloride is given to hyperthyroid patients or to those receiving thyroid medication, since cardiac arrhythmias may develop.

Nortriptyline hydrochloride may impair the mental and/or physical abilities required for the performance of hazardous tasks, such as operating machinery or driving a car; therefore, the patient should be warned accordingly.

Excessive consumption of alcohol in combination with nortriptyline therapy may have a potentiating effect, which may lead to the danger of increased suicidal attempts or overdosage, especially in patients with histories of emotional disturbances or suicidal ideation.

The concomitant administration of quinidine and nortriptyline may result in a significantly longer plasma half-life, higher AUC, and lower clearance of nortriptyline.

#### Serotonin Syndrome

The development of a potentially life-threatening serotonin syndrome has been reported with SNRIs and SSRIs, including nortriptyline hydrochloride, alone but particularly with concomitant use of other serotonergic drugs (including triptans, tricyclic antidepressants, fentanyl, lithium, tramadol, tryptophan, buspirone, and St. John's Wort) and with drugs that impair metabolism of serotonin (in particular, MAOIs, both those intended to treat psychiatric disorders and also others, such as linezolid and intravenous methylene blue).

Serotonin syndrome symptoms may include mental status changes (e.g., agitation, hallucinations, delirium, and coma), autonomic instability (e.g., tachycardia, labile blood pressure, dizziness, diaphoresis, flushing, hyperthermia), neuromuscular changes (e.g., tremor, rigidity, myoclonus, hyperreflexia, incoordination), seizures, and/or gastrointestinal symptoms (e.g., nausea, vomiting, diarrhea). Patients should be monitored for the emergence of serotonin syndrome.

The concomitant use of nortriptyline hydrochloride with MAOIs intended to treat psychiatric disorders is contraindicated. Nortriptyline hydrochloride should also not be started in a patient who is being treated with MAOIs such as linezolid or intravenous methylene blue. All reports with methylene blue that provided information on the route of administration involved intravenous administration in the dose

range of 1 mg/kg to 8 mg/kg. No reports involved the administration of methylene blue by other routes (such as oral tablets or local tissue injection) or at lower doses. There may be circumstances when it is necessary to initiate treatment with an MAOI such as linezolid or intravenous methylene blue in a patient taking nortriptyline hydrochloride. Nortriptyline hydrochloride should be discontinued before initiating treatment with the MAOI (see **CONTRAINDICATIONS** and **DOSAGE AND ADMINISTRATION**).

If concomitant use of nortriptyline hydrochloride with other serotonergic drugs, including triptans, tricyclic antidepressants, fentanyl, lithium, tramadol, buspirone, tryptophan, and St. John's Wort is clinically warranted, patients should be made aware of a potential increased risk for serotonin syndrome, particularly during treatment initiation and dose increases.

Treatment with nortriptyline hydrochloride and any concomitant serotonergic agents should be discontinued immediately if the above events occur and supportive symptomatic treatment should be initiated.

#### Use in Pregnancy

Safe use of nortriptyline hydrochloride during pregnancy and lactation has not been established; therefore, when the drug is administered to pregnant patients, nursing mothers, or women of childbearing potential, the potential benefits must be weighed against the possible hazards. Animal reproduction studies have yielded inconclusive results.

## PRECAUTIONS

#### **Information for Patients**

Prescribers or other health professionals should inform patients, their families, and their caregivers about the benefits and risks associated with treatment with nortriptyline hydrochloride capsules and should counsel them in their appropriate use. A patient Medication Guide about "Antidepressant Medicines, Depression and other Serious Mental Illnesses, and Suicidal Thoughts or Actions" is available for nortriptyline hydrochloride capsules. The prescriber or health professional should instruct patients, their families, and their caregivers to read the Medication Guide and should assist them in understanding its contents. Patients should be given the opportunity to discuss the contents of the Medication Guide and to obtain answers to any questions they may have. The complete text of the Medication Guide is reprinted at the end of this document.

Patients should be advised of the following issues and asked to alert their prescriber if these occur while taking nortriptyline hydrochloride capsules.

#### Clinical Worsening and Suicide Risk

Patients, their families, and their caregivers should be encouraged to be alert to the emergence of anxiety, agitation, panic attacks, insomnia, irritability, hostility, aggressiveness, impulsivity, akathisia (psychomotor restlessness), hypomania, mania, other unusual changes in behavior, worsening of depression, and suicidal ideation, especially early during antidepressant treatment and when the dose is adjusted up or down. Families and caregivers of patients should be advised to look for the emergence of such symptoms on a day-to-day basis, since changes may be abrupt. Such symptoms should be reported to the patient's prescriber or health professional, especially if they are severe, abrupt in onset, or were not part of the patient's presenting symptoms. Symptoms such as these may be associated with an increased risk for suicidal thinking and behavior and indicate a need for very close monitoring and possibly changes in the medication.

The use of nortriptyline hydrochloride in schizophrenic patients may result in an exacerbation of the psychosis or may activate latent schizophrenic symptoms. If the drug is given to overactive or agitated patients, increased anxiety and agitation may occur. In manic-depressive patients, nortriptyline hydrochloride may cause symptoms of the manic phase to emerge.

Troublesome patient hostility may be aroused by the use of nortriptyline hydrochloride. Epileptiform seizures may accompany its administration, as is true of other drugs of its class.

When it is essential, the drug may be administered with electroconvulsive therapy, although the hazards may be increased. Discontinue the drug for several days, if possible, prior to elective surgery.

The possibility of a suicidal attempt by a depressed patient remains after the initiation of treatment; in this regard, it is important that the least possible quantity of drug be dispensed at any given time.

Both elevation and lowering of blood sugar levels have been reported.

## **Drug Interactions**

Administration of reserpine during therapy with a tricyclic antidepressant has been shown to produce a "stimulating" effect in some depressed patients.

Close supervision and careful adjustment of the dosage are required when nortriptyline hydrochloride is used with other anticholinergic drugs and sympathomimetic drugs.

Concurrent administration of cimetidine and tricyclic antidepressants can produce clinically significant increases in the plasma concentrations of the tricyclic antidepressant. The patient should be informed that the response to alcohol may be exaggerated.

A case of significant hypoglycemia has been reported in a type II diabetic patient maintained on chlorpropamide (250 mg/day), after the addition of nortriptyline (125 mg/day).

## Drugs Metabolized by P450 2D6

The biochemical activity of the drug metabolizing isozyme cytochrome P450 2D6 (debrisoquin hydroxylase) is reduced in a subset of the Caucasian population (about 7% to 10% of Caucasians are so called "poor metabolizers"); reliable estimates of the prevalence of reduced P450 2D6 isozyme activity among Asian, African and other populations are not yet available. Poor metabolizers have higher than expected plasma concentrations of tricyclic antidepressants (TCAs) when given usual doses. Depending on the fraction of drug metabolized by P450 2D6, the increase in plasma concentration may be small, or quite large (8 fold increase in plasma AUC of the TCA).

In addition, certain drugs inhibit the activity of this isozyme and make normal metabolizers resemble poor metabolizers. An individual who is stable on a given dose of TCA may become abruptly toxic when given one of these inhibiting drugs as concomitant therapy. The drugs that inhibit cytochrome P450 2D6 include some that are not metabolized by the enzyme (quinidine; cimetidine) and many that are substrates for P450 2D6 (many other antidepressants, phenothiazines, and the Type 1C antiarrhythmics propafenone and flecainide). While all the selective serotonin reuptake inhibitors (SSRIs), e.g., fluoxetine, sertraline, and paroxetine, inhibit P450 2D6, they may vary in the extent of inhibition. The extent to which SSRI TCA interactions may pose clinical problems will depend on the degree of inhibition and the pharmacokinetics of the SSRI involved. Nevertheless, caution is indicated in the coadministration of TCAs with any of the SSRIs and also in switching from one class to the other. Of particular importance, sufficient time must elapse before initiating TCA treatment in a patient being withdrawn from fluoxetine, given the long half-life of the parent and active metabolite (at least 5 weeks may be necessary).

Concomitant use of tricyclic antidepressants with drugs that can inhibit cytochrome P450 2D6 may require lower doses than usually prescribed for either the tricyclic antidepressant or the other drug. Furthermore, whenever one of these other drugs is withdrawn from co-therapy, an increased dose of tricyclic antidepressant may be required. It is desirable to monitor TCA plasma levels whenever a TCA is going to be coadministered with another drug known to be an inhibitor of P450 2D6.

# Monoamine Oxidase Inhibitors (MAOIs)

(see **CONTRAINDICATIONS**, **WARNINGS**, and **DOSAGE AND ADMINISTRATION**).

#### Serotonergic Drugs

#### (see CONTRAINDICATIONS, WARNINGS, and DOSAGE AND ADMINISTRATION).

#### Pediatric Use

Safety and effectiveness in the pediatric population have not been established (see **BOX WARNING** and **WARNINGS**, **Clinical Worsening and Suicide Risk**). Anyone considering the use of nortriptyline hydrochloride capsules in a child or adolescent must balance the potential risks with the clinical need.

#### Geriatric Use

Clinical studies of nortriptyline hydrochloride did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. Other reported clinical experience indicates that, as with other tricyclic antidepressants, hepatic adverse events (characterized mainly by jaundice and elevated liver enzymes) are observed very rarely in geriatric patients and deaths associated with cholestatic liver damage have been reported in isolated instances. Cardiovascular function, particularly arrhythmias and fluctuations in blood pressure, should be monitored. There have also been reports of confusional states following tricyclic antidepressant administration in the elderly. Higher plasma concentrations of the active nortriptyline metabolite, 10-hydroxynortriptyline, have also been reported in elderly patients. As with other tricyclic antidepressants, dose selection for an elderly patient should usually be limited to the smallest effective total daily dose (see **DOSAGE AND ADMINISTRATION**).

#### **ADVERSE REACTIONS**

**Note** - Included in the following list are a few adverse reactions that have not been reported with this specific drug. However, the pharmacologic similarities among the tricyclic antidepressant drugs require that each of the reactions be considered when nortriptyline is administered.

*Cardiovascular* - Hypotension, hypertension, tachycardia, palpitation, myocardial infarction, arrhythmias, heart block, stroke.

*Psychiatric* - Confusional states (especially in the elderly) with hallucinations, disorientation, delusions; anxiety, restlessness, agitation; insomnia, panic, nightmares; hypomania; exacerbation of psychosis.

*Neurologic* - Numbness, tingling, paresthesias of extremities; incoordination, ataxia, tremors; peripheral neuropathy; extrapyramidal symptoms; seizures, alteration in EEG patterns; tinnitus.

**Anticholinergic** - Dry mouth and, rarely, associated sublingual adenitis; blurred vision, disturbance of accommodation, mydriasis; constipation, paralytic ileus; urinary retention, delayed micturition, dilation of the urinary tract.

*Allergic* - Skin rash, petechiae, urticaria, itching, photosensitization (avoid excessive exposure to sunlight); edema (general or of face and tongue), drug fever, cross-sensitivity with other tricyclic drugs.

*Hematologic* - Bone marrow depression, including agranulocytosis; eosinophilia; purpura; thrombocytopenia.

*Gastrointestinal* - Nausea and vomiting, anorexia, epigastric distress, diarrhea, peculiar taste, stomatitis, abdominal cramps, black-tongue.

*Endocrine* - Gynecomastia in the male, breast enlargement and galactorrhea in the female; increased or decreased libido, impotence; testicular swelling; elevation or depression of blood sugar levels; syndrome of inappropriate ADH (antidiuretic hormone) secretion.

**Other** - Jaundice (simulating obstructive), altered liver function; weight gain or loss; perspiration; flushing; urinary frequency, nocturia; drowsiness, dizziness, weakness, fatigue; headache; parotid swelling; alopecia.

*Withdrawal Symptoms* - Though these are not indicative of addiction, abrupt cessation of treatment after prolonged therapy may produce nausea, headache, and malaise.

## OVERDOSAGE

Deaths may occur from overdosage with this class of drugs. Multiple drug ingestion (including alcohol) is common in deliberate tricyclic antidepressant overdose. As the management is complex and changing, it is recommended that the physician contact a poison control center for current information on treatment. Signs and symptoms of toxicity develop rapidly after tricyclic antidepressant overdose, therefore, hospital monitoring is required as soon as possible.

#### **Manifes** tations

Critical manifestations of overdose include: cardiac dysrhythmias, severe hypotension, shock, congestive heart failure, pulmonary edema, convulsions, and CNS depression, including coma. Changes in the electrocardiogram, particularly in QRS axis or width, are clinically significant indicators of tricyclic antidepressant toxicity.

Other signs of overdose may include: confusion, restlessness, disturbed concentration, transient visual hallucinations, dilated pupils, agitation, hyperactive reflexes, stupor, drowsiness, muscle rigidity, vomiting, hypothermia, hyperpyrexia, or any of the acute symptoms listed under **ADVERSE REACTIONS**. There have been reports of patients recovering from nortriptyline overdoses of up to 525 mg.

#### Management

#### General

Obtain an ECG and immediately initiate cardiac monitoring. Protect the patient's airway, establish an intravenous line and initiate gastric decontamination. A minimum of six hours of observation with cardiac monitoring and observation for signs of CNS or respiratory depression, hypotension, cardiac dysrhythmias and/or conduction blocks, and seizures is necessary. If signs of toxicity occur at any time during this period, extended monitoring is required. There are case reports of patients succumbing to fatal dysrhythmias late after overdose; these patients had clinical evidence of significant poisoning prior to death and most received inadequate gastrointestinal decontamination. Monitoring of plasma drug levels should not guide management of the patient.

#### Gas trointes tinal Decontamination

All patients suspected of tricyclic antidepressant overdose should receive gastrointestinal decontamination. This should include large volume gastric lavage followed by activated charcoal. If consciousness is impaired, the airway should be secured prior to lavage. EMESIS IS CONTRAINDICATED.

## **Cardiovas cular**

A maximal limb-lead QRS duration of  $\geq 0.10$  seconds may be the best indication of the severity of the overdose. Intravenous sodium bicarbonate should be used to maintain the serum pH in the range of 7.45 to 7.55. If the pH response is inadequate, hyperventilation may also be used. Concomitant use of hyperventilation and sodium bicarbonate should be done with extreme caution, with frequent pH monitoring. A pH > 7.60 or a pCO<sub>2</sub> < 20 mmHg is undesirable. Dysrhythmias unresponsive to sodium bicarbonate therapy/hyperventilation may respond to lidocaine, bretylium or phenytoin. Type 1A and 1C antiarrhythmics are generally contraindicated (e.g., quinidine, disopyramide, and procainamide). In rare instances, hemoperfusion may be beneficial in acute refractory cardiovascular instability in patients with acute toxicity. However, hemodialysis, peritoneal dialysis, exchange transfusions, and forced diuresis generally have been reported as ineffective in tricyclic antidepressant poisoning.

## CNS

In patients with CNS depression, early intubation is advised because of the potential for abrupt deterioration. Seizures should be controlled with benzodiazepines, or if these are ineffective, other anticonvulsants (e.g., phenobarbital, phenytoin). Physostigmine is not recommended except to treat life-threatening symptoms that have been unresponsive to other therapies, and then only in consultation with a poison control center.

#### Psychiatric Follow-up

Since overdosage is often deliberate, patients may attempt suicide by other means during the recovery phase. Psychiatric referral may be appropriate.

## **Pediatric Management**

The principles of management of child and adult overdosages are similar. It is strongly recommended that the physician contact the local poison control center for specific pediatric treatment.

## DOSAGE AND ADMINISTRATION

Nortriptyline hydrochloride is not recommended for children.

Nortriptyline hydrochloride is administered orally in the form of capsules. Lower than usual dosages are recommended for elderly patients and adolescents. Lower dosages are also recommended for outpatients than for hospitalized patients who will be under close supervision. The physician should initiate dosage at a low level and increase it gradually, noting carefully the clinical response and any evidence of intolerance. Following remission, maintenance medication may be required for a longer period of time at the lowest dose that will maintain remission.

If a patient develops minor side effects, the dosage should be reduced. The drug should be discontinued promptly if adverse effects of a serious nature or allergic manifestations occur.

## Usual Adult Dose

25 mg three or four times daily; dosage should begin at a low level and be increased as required. As an alternate regimen, the total daily dosage may be given once a day. When doses above 100 mg daily are administered, plasma levels of nortriptyline should be monitored and maintained in the optimum range of 50 to 150 mg/mL. Doses above 150 mg/day are not recommended.

## **Elderly and Adoles cent Patients**

30 to 50 mg/day, in divided doses, or the total daily dosage may be given once a day.

# Switching a Patient To or From a Monoamine Oxidase Inhibitor (MAOI) Intended to Treat Psychiatric Disorders

At least 14 days should elapse between discontinuation of an MAOI intended to treat psychiatric disorders and initiation of therapy with nortriptyline hydrochloride. Conversely, at least 14 days should be allowed after stopping nortriptyline hydrochloride before starting an MAOI intended to treat psychiatric disorders (see **CONTRAINDICATIONS**).

# Use of Nortriptyline Hydrochloride With Other MAOIs, Such as Linezolid or Methylene Blue

Do not start nortriptyline hydrochloride in a patient who is being treated with linezolid or intravenous methylene blue because there is increased risk of serotonin syndrome. In a patient who requires more urgent treatment of a psychiatric condition, other interventions, including hospitalization, should be considered (see **CONTRAINDICATIONS**).

In some cases, a patient already receiving nortriptyline hydrochloride therapy may require urgent

treatment with linezolid or intravenous methylene blue. If acceptable alternatives to linezolid or intravenous methylene blue treatment are not available and the potential benefits of linezolid or intravenous methylene blue treatment are judged to outweigh the risks of serotonin syndrome in a particular patient, nortriptyline hydrochloride should be stopped promptly, and linezolid or intravenous methylene blue can be administered. The patient should be monitored for symptoms of serotonin syndrome for two weeks or until 24 hours after the last dose of linezolid or intravenous methylene blue, whichever comes first. Therapy with nortriptyline hydrochloride may be resumed 24 hours after the last dose of linezolid or intravenous methylene blue,

The risk of administering methylene blue by non-intravenous routes (such as oral tablets or by local injection) or in intravenous doses much lower than 1 mg/kg with nortriptyline hydrochloride is unclear. The clinician should, nevertheless, be aware of the possibility of emergent symptoms of serotonin syndrome with such use (see **WARNINGS**).

## HOW SUPPLIED

Nortriptyline hydrochloride capsules USP, equivalent to 25 mg, 50 mg or 75 mg base, are available as follows:

Nortriptyline hydrochloride capsules USP, 25 mg: #4 capsules with a white body and an orange cap, imprinted "TEVA"-"TEVA" on the cap and "25mg"-"0811" on the body, in bottles of 30, 60 and 120.

Nortriptyline hydrochloride capsules USP, 50 mg: #1 capsules with a white body and a white cap, imprinted "TEVA" on the cap and "50mg"-"0812" on the body, in bottles of 30 and 60.

Nortriptyline hydrochloride capsules USP, 75 mg: #1 capsules with an orange body and an orange cap, imprinted "TEVA" on the cap and "75mg"-"0813" on the body, in bottles of 30.

Store at 20° to 25°C (68° to 77°F) [See USP Controlled Room Temperature].

Dispense contents in a tight container as defined in the USP, with a child-resistant closure (as required).

# **TEVA PHARMACEUTICALS USA**

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## **MEDICATION GUIDE**

## Nortriptyline Hydrochloride Capsules USP

# Antidepressant Medicines, Depression and other Serious Mental Illnesses, and Suicidal Thoughts or Actions

#### **Rx only**

Read the Medication Guide that comes with your or your family member's antidepressant medicine. This Medication Guide is only about the risk of suicidal thoughts and actions with antidepressant medicines. **Talk to your, or your family member's, healthcare provider about:** 

- all risks and benefits of treatment with antidepressant medicines
- all treatment choices for depression or other serious mental illness

# What is the most important information I should know about antidepressant medicines, depression and other serious mental illnesses, and suicidal thoughts or actions?

1. Antidepressant medicines may increase suicidal thoughts or actions in some children,

#### teenagers, and young adults within the first few months of treatment.

**2.Depression and other serious mental illnesses are the most important causes of suicidal thoughts and actions. Some people may have a particularly high risk of having suicidal thoughts or actions.** These include people who have (or have a family history of) bipolar illness (also called manic-depressive illness) or suicidal thoughts or actions.

# 3. How can I watch for and try to prevent suicidal thoughts and actions in myself or a family member?

- Pay close attention to any changes, especially sudden changes, in mood, behaviors, thoughts, or feelings. This is very important when an antidepressant medicine is started or when the dose is changed.
- Call the healthcare provider right away to report new or sudden changes in mood, behavior, thoughts, or feelings.
- Keep all follow-up visits with the healthcare provider as scheduled. Call the healthcare provider between visits as needed, especially if you have concerns about symptoms.

# Call a healthcare provider right away if you or your family member has any of the following symptoms, especially if they are new, worse, or worry you:

- thoughts about suicide or dying
- attempts to commit suicide
- new or worse depression
- new or worse anxiety
- feeling very agitated or restless
- panic attacks
- trouble sleeping (insomnia)
- new or worse irritability
- acting aggressive, being angry, or violent
- acting on dangerous impulses
- an extreme increase in activity and talking (mania)
- other unusual changes in behavior or mood

# Who should not take nortriptyline hydrochloride?

Do not take nortriptyline hydrochloride if you:

• take a monoamine oxidase inhibitor (MAOI). Ask your healthcare provider or pharmacist if you are not sure if you take an MAOI, including the antibiotic linezolid.

° Do not take an MAOI within 2 weeks of stopping nortriptyline hydrochloride unless

directed to do so by your physician.

° Do not start nortriptyline hydrochloride if you stopped taking an MAOI in the last 2 weeks

unless directed to do so by your physician.

# What else do I need to know about antidepressant medicines?

- **Never stop an antidepressant medicine without first talking to a healthcare provider.** Stopping an antidepressant medicine suddenly can cause other symptoms.
- Antidepressants are medicines used to treat depression and other illnesses. It is important to discuss all the risks of treating depression and also the risks of not treating it. Patients and their families or other caregivers should discuss all treatment choices with the healthcare provider, not just the use of antidepressants.
- **Antidepressant medicines have other side effects.** Talk to the healthcare provider about the side effects of the medicine prescribed for you or your family member.
- Antidepressant medicines can interact with other medicines. Know all of the medicines that you

or your family member takes. Keep a list of all medicines to show the healthcare provider. Do not start new medicines without first checking with your healthcare provider.

• Not all antidepressant medicines prescribed for children are FDA approved for use in children. Talk to your child's healthcare provider for more information.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

This Medication Guide has been approved by the U.S. Food and Drug Administration.

## **TEVA PHARMACEUTICALS USA**

Sellersville, PA 18960

Rev. F 11/2012

#### PRINCIPAL DISPLAY PANEL

Packaged and Distributed by:	AIDAREX PHA	RMACEUTICALS LLC.	NORTRIPTYLINE 25 MG
	NORTE		NDC: 33261-0348-30 RX QLS0000 30
		()	NORTRIPTYLINE 25 MG 👳
ROOM T	25 MG	ACTIVE INGREDIENTS:	NDC: 33261-0348-30
	30 CAPS	OR AN GE AND WHITE OFLONG CAPSULE W/ TEVA 25M G ON ONE SIDE & TEVA 0811 ON REVERSE	NORTRIPTYLINE 25 MG
	VERIC FOR : PAMELOF	4	RXQLS0000 30
NDC	C: 33261-0348-30	AKEEVERYHOURSTIMES A DAY	
E.		OME <u>CADA HORAS</u> VECES AL DIA CALS USA SELLERSVILLE, PA 18960	NORTRIPTYLINE 25 MG N DC: 33261-0348-30 RX QLS0000 30

Nortriptyline HCl Capsules 25 mg 30s Label Text

NDC 33261-0348-30

NORTRIPTYLINE

#### HYDROCHLORIDE

#### **Capsules USP**

25 mg\*

PHARMACIST: Dispense the

accompanying Medication Guide

to each patient.

Rx only

**30 CAPSULES** 

PRINCIPAL DISPLAY PANEL

u (Packag O 8Distribu T d	ed and AIDAREX PHA	RMACEUTICALS LLC.		NORTRIPTYLINE	50 MG	PATIENT	
SSOC(SC	NORTE	RIPTYLINE	0000	NDC: 33261-0692-30 RX QLS0000	30	TAT	
		EACH CAPSULE CONTAINS THE FOLLOWING	3	NORTRIPTYLINE	50 MG	DOT	
	50 MG	ACTIVE INGREDIENTS: NORTRIPTYLINE HYDRO CHLO RIDE	a	NDC: 33261-0692-30 RX QLS0000	0 30	I	
	30 CAPS	WHITE OBLONG CAPSULE W/TEVA 50MG 0812 ON ONE SIDE	Ê	NORTRIPTYLINE NDC: 33261-0692-34	50 MG	CHART	
ATOA	GENERIC FOR : PAMELOR			RX QLS0000	30		
CAUTION: FEDERAL PRESCAIPTION. SE CHILDREN. STORE/	100201-0002-00	KEEVERYHOURSTIMES MECADAHORASVECES SUSA SELLERSVILLE, PA 18960		NORTRIPTYLINE NDC: 33261-0692-3 RX QLS0000	50 MG 0 30		PELI.IERE

Nortriptyline HCl Capsules 50 mg 30s Label Text

NDC 33261-0692-30

## NORTRIPTYLINE

# HYDROCHLORIDE

**Capsules USP** 

50 mg\*

PHARMACIST: Dispense the accompanying

Medication Guide to each patient.

Rx only

**30 CAPSULES** 

# PRINCIPAL DISPLAY PANEL

B B Packaged an		HARMACEUTICALS LLC.	NORTRIPTYLINE 75 mg	LUE
COUT OF REAC	NORT		RX QLS0000 30	TATENT TATENT
	75 mg	ACTIVE INGREDIENTS:	NDC: 33261-0804-30 RX QLS0000 30	TOG
	30 CAPS	ORANGE OBLONG CAPSULE W/ TEVA 75 MG 0813 ON ONE SIDE	NORTRIPTYLINE <b>75 mg</b> NDC: 33261-0804-30 RXGLS0000 30	CHART
Шш	IDC: 33261-0804-30	TAKEEVERYHOURSTIMES ADAY		PEEL
	TEVA PHARMACEUTICAL	TOMECADAHORASVECES AL DIA S USA SELLERSVILLE, PA 18960	NORTRIPTYLINE 75 mg NDC: 33261-0804-30 RX QLS0000 30	JL III.RE TTE

# Nortriptyline HCl Capsules 75 mg 30s Label Text

NDC 33261-0804-30

NORTRIPTYLINE

## HYDROCHLORIDE

Capsules USP

# 75 mg\*

PHARMACIST: Dispense the accompanying

Medication Guide to each patient.

#### Rx only

**30 CAPSULES** 

Product Information				
Product T ype	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:33261-348(1	NDC:0093-0811)
Route of Administration	ORAL			
Active Ingredient/Active	Moiety			
	Ingredient Name		Basis of Strengtl	Strengt
NORTRIPTYLINE HYDROCHLO UNII:BL03SY4LXB)	RIDE (UNII: 00FN6IH15D) (NORTRIP	TYLINE -	NORTRIPTYLIN	NE 25 mg
Inactive Ingredients				
inactive ingredients	Ingredient Name			Strength
STARCH, CORN (UNII: 08232NY	C C			ourengen
D&C YELLOW NO. 10 (UNII: 355				
ALUMINUM OXIDE (UNII: LMI26				
FD&C BLUE NO. 1 (UNII: H3R47k	(3TBD)			
FD&C BLUE NO. 2 (UNII: L06K8	R7DQK)			
FD&C RED NO. 40 (UNII: WZB91	27XOA)			
GELATIN (UNII: 2G86QN327L)				
FERROSOFERRIC OXIDE (UNII:	XM0 M8 7F357)			
METHYLPARABEN (UNII: A218C)	7HI9T)			
PROPYLENE GLYCOL (UNII: 6D	C9Q167V3)			
PROPYLPARABEN (UNII: Z8IX25	SC10H)			
<b>SHELLAC</b> (UNII: 46 N107B710)				
DIMETHICO NE (UNII: 92RU3N3Y				
SODIUM LAURYL SULFATE (UI	,			
TITANIUM DIO XIDE (UNII: 15FIX	9 V2 ID)			

<b>Product Ch</b>	aracteris	stics							
Color	WHIT	E, ORANGE		Score		no score			
Shape	CAPS	ULE		Size		14mm			
Flavor				Imprint Code		TEVA;TEV	TEVA;TEVA;25mg;0811		
Contains									
Packaging									
# Item	Code	Pac	kage Descript	ion I	Marketi	ing Start Date	Μ	arketing <b>E</b>	nd Date
1 NDC:33261-3	348-02	120 in 1 B	OTTLE, PLASTIC	2					
2 NDC:33261-3	348-30	30 in 1 BC	TTLE, PLASTIC						
<b>3</b> NDC:33261-3	348-60	60 in 1 BC	OTTLE, PLASTIC						
Marketin	g Infor	mation							
Marketing C	<b>-</b>		n Number er N	Jongraph C:	tation	Markating St	nt Data	Mankatin	a End Data
-			on Number or N	lonograph Cl	graph Citation Marketing Start			t Date Marketing End	
ANDA	P	ANDA074132				04/06/1995			
			ROCHLOF	RIDE					
nortriptyline h	ydrochlori	ide capsule	ROCHLOF	RIDE					
nortriptyline h Product In	ydrochlori f <b>ormatior</b>	ide capsule	ROCHLOF		Item (	Code (Source)	NDC:3320	51-692(NDC	:0093-0812)
nortriptyline h Product In	ydrochlori formatior e	ide capsule 1			Item(	Code (Source)	NDC:3320	51-692(NDC	:0093-0812)
nortriptyline h Product Inf Product Typ	ydrochlori formatior e	ide capsule 1	HUMAN PRESC		Item(	Code (Source)	NDC:3320	51-692(NDC)	:0093-0812)
nortriptyline h Product Inf Product Typ	ydrochlori f <b>ormatior</b> e ninistratior	ide capsule 1	HUMAN PRESC ORAL		Item	Code (Source)			:0093-0812)
nortriptyline h Product Inf Product Typ Route of Adm	ydrochlori f <b>ormatior</b> e ninistratior	ide capsule 1 1 ctive Moie	HUMAN PRESC ORAL	RIPTION DRUG	Item (	Code (Source)	B	51-692(NDC asis of trength	:0093-0812) Strengt
nortriptyline h Product Inf Product Type Route of Adm Active Ingr	ydrochlori formatior e ninistratior e die nt/Ad	ide capsule 1 1 ctive Moie	HUMAN PRESC ORAL	RIPTION DRUG			B	asis of	
nortriptyline h Product Inf Product Typ Route of Adm Active Ingr NORTRIPTYL	ydrochlori formatior e ninistratior e die nt/Ad	ide capsule 1 1 ctive Moie	HUMAN PRESC ORAL ety Ingredient Na	RIPTION DRUG			B	asis of trength	Strengt
nortriptyline h Product Inf Product Typ Route of Adm Active Ingr NORTRIPTYL	ydrochlori formation e ninistration e die nt/Ao INE HYDRO LXB)	ide capsule 1 1 ctive Moie	HUMAN PRESC ORAL ety Ingredient Na E (UNII: 00FN6IH	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	asis of trength RIPTYLINE	Strengt 50 mg
ortriptyline h Product Inf Product Type Route of Adm Active Ingr NORTRIPTYL UNII:BL03SY41	ydrochlori formation e ninistration e die nt/Ao INE HYDRO LXB)	ide capsule 1 1 ctive Moie	HUMAN PRESC ORAL ety Ingredient Na	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	Gasis of trength RIPTYLINE	Strengt
Product Ind Product Ind Product Type Route of Adm Active Ingr NORTRIPTYL UNII:BL03SY41 Inactive Ing STARCH, COF	ydrochlori formatior e ninistratior e die nt/Ad INE HYDRC _XB) gre die nts	ide capsule 1 1 Ctive Moio OCHLORIDE	HUMAN PRESC ORAL ety Ingredient Na E (UNII: 00FN6 IH Ingredient	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	Gasis of trength RIPTYLINE	Strengt 50 mg
Product Inf Product Inf Product Type Route of Adm Active Ingr NORTRIPTYL UNII:BL03SY41 Inactive Ing STARCH, COF D&C YELLOV	ydrochlori formation e ninistration e die nt/Ad e die nt/Ad INE HYDRO LXB) gre die nts gre die nts RN (UNII: 08 V NO. 10 (U	ide capsule 1 1 1 1 1 1 1 1 1 1 1 1 1	HUMAN PRESC ORAL ety Ingredient Na E (UNII: 00FN6IH Ingredient	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	Gasis of trength RIPTYLINE	Strengt 50 mg
Product Inf Product Inf Product Type Route of Adm Active Ingr NORTRIPTYL UNII:BL03SY4I Inactive Ing STARCH, COF D&C YELLOV ALUMINUM O	ydrochlori formation e ninistration e die nt/Ad e die nt/Ad iNE HYDRO XB) gre die nts gre die nts N (UNII: 08 v NO. 10 (U XIDE (UNII:	ide capsule 1 1 1 1 1 1 1 1 1 1 1 1 1	HUMAN PRESC ORAL ety Ingredient Na E (UNII: 00FN6IH Ingredien SQ3G) 3)	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	Gasis of trength RIPTYLINE	Strengt 50 mg
nortriptyline h Product Inf Product Type Route of Adm Active Ingr NORTRIPTYL UNII:BL03SY41 Inactive Ing STARCH, COF D&C YELLOV	ydrochlori formation e ninistration e die nt/Ad e die nt/Ad iNE HYDRO XB) gre die nts gre die nts N (UNII: 08 v NO. 10 (U XIDE (UNII:	ide capsule 1 1 1 1 1 1 1 1 1 1 1 1 1	HUMAN PRESC ORAL ety Ingredient Na E (UNII: 00FN6IH Ingredien SQ3G) 3)	RIPTION DRUG I <b>me</b> 15D) (NORTRIP			B	Gasis of trength RIPTYLINE	Strengt 50 mg

 FD&C RED NO. 40 (UNII: WZB9127XOA)

 GELATIN (UNII: 2G86QN327L)

 FERROSOFERRIC OXIDE (UNII: XM0M87F357)

METHYLPARABEN (UNII: A2I8C7HI9T) PROPYLENE GLYCOL (UNII: 6DC9Q167V3) PROPYLPARABEN (UNII: Z8IX2SC10H)

SHELLAC (UNII: 46 N107	B71O)							
DIMETHICO NE (UNII: 92	2RU3N3Y1O)							
SODIUM LAURYL SULI	F <b>ATE</b> (UNII: 368GB5141J)							
TITANIUM DIO XIDE (U	NII: 15FIX9V2JP)							
<b>Product Character</b>	istics							
Color	Color WHITE Score no score							
Shape	CAPSULE	Size			19 mm			
Flavor		Imprint Cod	le		TEVA;50m	g;0812	2	
Contains								
Packaging								
# Item Code	Package Desc	ription	Marketi	ng Start	Date	M	arketing End Date	
<b>1</b> NDC:33261-692-30	30 in 1 BOTTLE, PLA	STIC						
<b>2</b> NDC:33261-692-60	60 in 1 BOTTLE, PLA	STIC						
Marketing Info	rmation							
			h Citatian	Maulta	ing Casul I	Data	Maulating End Data	
Marketing Category	Application Number	or Monogra	on Citation	Marketing Start Date			Marketing End Date	
ANDA	ANDA074132			04/05/19	95			
NORTRIPTYLI	NE HYDROCHL	ORIDE						
<b>NORTRIPTYLI</b> nortriptyline hydrochlo		ORIDE						

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:33261-804(NDC:0093-0813)
Route of Administration	ORAL		

	D,	asis of	
Ingredient Name		rength	Strengt
NORTRIPTYLINE HYDROCHLORIDE (UNII: 00FN6IH15D) (NORTRIPTYLINE - UNII:BL03SY4LXB)	NORTRI	PTYLINE	75 mg
Inactive Ingredients			
Ingredient Name		Strei	ıgth
Ingreutent Name			0
STARCH, CORN (UNII: 08232NY3SJ)			8
STARCH, CORN (UNII: 08232NY3SJ)			0
STARCH, CORN (UNII: 08232NY3SJ) D&C YELLOW NO. 10 (UNII: 35SW5USQ3G)			0
STARCH, CORN (UNII: 08232NY3SJ) D&C YELLOW NO. 10 (UNII: 35SW5USQ3G) ALUMINUM OXIDE (UNII: LMI2606933)			0
			0

GELATIN (UNII: 2G86QN	I327L)						
FERROSOFERRIC OXID	<b>DE</b> (UNII: XM0 M8 7F357)						
METHYLPARABEN (UNI	I: A2I8C7HI9T)						
PROPYLENE GLYCOL (	(UNII: 6DC9Q167V3)						
PROPYLPARABEN (UNI	I: Z8IX2SC1OH)						
SHELLAC (UNII: 46 N107	B71O)						
DIMETHICONE (UNII: 92	RU3N3Y1O)						
SODIUM LAURYL SULF	<b>ATE</b> (UNII: 368GB5141J)						
TITANIUM DIO XIDE (UN	NII: 15FIX9V2JP)						
FD&C YELLOW NO.6 (	UNII: H77VEI93A8)						
<b>Product Characteri</b>	istics						
Color	ORANGE	Score			no score		
Shape	CAPSULE	Size			19 mm		
Flavor		Imprint Cod	le		TEVA;75n	ng;0813	5
Contains							
Packaging							
# Item Code	Package Descr	ription	Marketi	ng Start i	Date	Ma	arketing End Date
1 NDC:33261-804-30	30 in 1 BOTTLE, PLAS	TIC					
Maultating Infor	mation						
Marketing Infor							
Marketing Category	Application Number	or Monograp	h Citation		ing Start	Date	Marketing End Date
ANDA	ANDA074132			04/05/199	95		

Labeler - Aidarex Pharmaceuticals LLC (801503249)

Revised: 1/2014

Aidarex Pharmaceuticals LLC