Rischotic[®] OD

Risperidone

FORMS AND PRESENTATION
Rischotic® OD 0.5: Orally disintegrating tablets; Box of 30.
Rischotic® OD 1: Orally disintegrating tablets: Box of 30.
Rischotic® OD 2: Orally disintegrating tablets: Box of 30.
Rischotic® OD 2: Orally disintegrating tablets: Box of 30.
Rischotic® OD 4: Orally disintegrating tablets: Box of 30.
Rischotic® OD 4: Orally disintegrating tablets: Box of 30.
Rischotic® OD 5: Each orally disintegrating tablets: Box of 30.
Rischotic® OD 5: Each orally disintegrating tablet contains Risperidone 1mg.
Rischotic® OD 5: Each orally disintegrating tablet contains Risperidone 2mg.
Rischotic® OD 2: Each orally disintegrating tablet contains Risperidone 3mg.
Rischotic® OD 3: Each orally disintegrating tablet contains Risperidone 4mg.
Excipients: magnesium aluminometasilicate, aspartame, colloidal silicon dioxide,
hydroxyproyl cellulose, sodium steary fumarate, methacrylate cooplymer, tale, sodium
lauryl sulfate, assulfame potassium, mannitol, sodium chloride, peppermint flavor,
yellow iron oxide (Rischotic® OD 4).
PHARNACOLOGICAL PROPERTIES
Pharmacodynamic properties:
ATIC code: NOAX.NOS.
Risperidone is a selective commaninergie antagonist with unique properties. It has a high
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ATC čode: NO5AX08. Klisperidone is a selective monoaminergic antagonist with unique properties. It has a high affinity for serotoninergic 5-HT2 and dopaminergic D2 receptors. Risperidone binds also to alphal-adrenergic receptors, and, with lower affinity, to H1-histaminergic and alpha2-adrenergic receptors. Risperidone has no affinity for cholinergic receptors. Although Risperidone is a potent D2 antagonist, which is considered to improve the positive symptoms of schizophrenia, it causes less depression of motor activity and dopamine antagonism may reduce extrapyramidal side effect liability and extend the therapeutic activity to the negative and affective symptoms of schizophrenia. *Pharmacokinetic properties* Absorption

Pharmacokinetic properties Absorption Risperidone is completely absorbed after oral administration, reaching peak plasma concentrations within 1 to 2 hours. The relative oral bioavailability of Risperidone from a tablet is 94% compared with a solution. <u>Distribution</u> Risperidone is rapidly distributed. The plasma protein binding of Risperidone is 90%, that of 9-hydroxy-risperidone is 77%. Risperidone plasma concentrations are dose-proportional within the therapeutic dose-range.

The state of the short-term treatment (up to 6 weeks) of persistent augments in state to 1 superscription of the short-term treatment (up to 6 weeks) of persistent augments and the big to 1 superscription of the short-term treatment (up to 6 weeks) of persistent augments and the big to 1 superscription of the short-term treatment (up to 6 weeks) of persistent augments and the big to 1 superscription of the short-term treatment (up to 6 weeks) of persistent augments in superscription of the short-term treatment (up to 6 weeks) of persistent augments in superscription of the short-term treatment (up to 6 weeks) of persistent augments in superscription augments in superscription augments and the big augments augments and the short-term treatment (up to 6 weeks) of persistent augments in superscription augments in superscription augments in superscription augments in superscription augments and the short-term treatment (up to 6 weeks) of persistent augments in superscription augments in the superscription augments in the superscription augments in superscription augments in superscription augments augments in the superscription augments augm

Por the treatment of schuzophrenia.
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 For the schuzephrenia of moderate to severe manic episodes associated with bipolar for the schuzephrene to severe Alzheimer's dementia unresponsive to non-pharmacological approaches and when there is a risk of harm to self or others.
 For the short-term symptomatic treatment (up to 6 weeks) of persistent aggression in patients with moderate to severe Alzheimer's dementia unresponsive to non-pharmacological approaches and when there is a risk of harm to self or others.
 For the short-term symptomatic treatment (up to 6 weeks) of persistent aggression in conduct disorder in children from the age of 5 years and adolescents with sub average intellectual functioning or mental netardation diagnosed according to DSM-IV criteria. CONTRAINDICATIONS
 Hypersensitivity to Risperidone or to any of the excipients.
 PRECAUTONS
 Elderly patients with dementia
 Overall mortality: Elderly patients with dementia treated with atypical antipsychotics, including Risperidone.
 Cerebrovascular events, including stroke, in elderly patients with dementia-related psychosis: Risperidone® OD is not approved for use in patients with demeta-related psychosis.
 Tardive dyskinesia: Consider discontinuing Risperidone® OD if clinically indicated.
 Metabolic Changes: Atypical antipsychotic drugs have been associated with metabolic changes that may increase cardiovascular' cerebroascular'.
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administration. Portex and the provided of the

Selzültes: Use cautous, in present conditions that lower the seizure threshold. Excipients
 The orodispersible tablets contain aspartame. Aspartame is a source of phenylalanine which may be harmful for people with phenylketonuria.
 PREGNANCY AND LACTATION
 There are no adequate data from the use of Risperidone in pregnant women. Neonates exposed to antipsychotics during the third trimster of pregnancy are at risk of adverse reactions including extrapyramidal and/or withdrawal symptoms that may vary in severity and duration. There have been reports of agitation, hypertonia, hypotonia, tremor, somnolence, respiratory distress, or feeding disorder. Risperidone was not teratogenic in animal studies but other types of reproductive toxicity were seen. Risperidone should not be used during pregnancy unless clearly necessary. If discontinuation during pregnancy is necessary, it should not be done abrupty.
 It has been demonstrated that Risperidone and 9-hydroxy-risperidone are excreted in human breast milk in small quantities. There are no data available on adverse reactions in breast-feeding infants. Therefore, the advantage of breastfeeding should be weighed against the potential risks for the child.
 DRUG INTERACTIONS.
 As with other antipsychotics, caution is advised when prescribing Risperidone with available on to routous to routous the OT interval, e.g., class Ia antiartifythinics and source in the readvant is more when to morigone the OT interval, e.g., class Ia antiartifythinics and source and antipsychotics.

against the potential risks for the child. **DRUG INTERACTIONS** As with other antipsychotics, caution is advised when prescribing Risperidone with medicinal products known to prolong the QT interval, e.g., class I aniaritythninics and class III antiaritythmics tricyclic antidepressant, tetracyclic antidepressants, some antihistaminics, other antipsychotics, some antimalaritals (i.e., chinice and methoquine), and with medicines causing electrolyte imbalance (hypokalemia, hypomagnesiemia), bradycardia, or those which inhibit the hepatic metabolism of Risperidone. Potential for Risperidone to affect other medicinal products - Risperidone should be used with caution in combination with other centrally-acting substances notably including alcohol, opiates, antihistamines and benzodiazepines due to the increased risk of sedation. - Risperidone may antagonize the effect of levodopa and other dopamine agonists. If this combination is deemed necessary, particularly in end-stage Parkinson's disease, the lowest effective dose of each treatment should be prescribed. - Clinically significant hypotension has been observed postmarketing with concomitant use of Risperidone and antihypertensive treatment. Potential for other medicinal products to affect Risperidone - Carbamazepine has been shown to decrease the plasma concentrations of the active antipsychotic fraction of Risperidone. Similar effects may be observed with e.g. rifampicin, phenytoin and phenobarbital - Fluoxetine and paroxeture, CYP 2D6 inhibitors, increase the plasma concentrations of Risperidone, but less so of the active antipsychotic fraction. It is expected that other CYP 2D6 inhibitors, such as quinidine, may affect the plasma concentrations of Risperidone a similar way. - Veranamil, an inbihitor of CYP 3A4 and P-on increases the plasma concentration of

2D0 initiotors, such as quintume, may anect the plasma concentrations of Risperdone in a similar way. - Verapamil, an inhibitor of CYP 3A4 and P-gp, increases the plasma concentration of Risperidone. - Phenothiazines, tricyclic antidepressants, and some beta-blockers may increase the plasma concentrations of Risperidone but not those of the active antipsychotic fraction. Cimetidine and ranitidine increase the bioavailability of Risperidone, but only marginally that of the active antipsychotic fraction. Erythornycin, a CYP 3A4 inhibitor, does not change the pharmacokinetics of Risperidone and the active antipsychotic fraction.

fraction. - Concomitant use of oral Risperidone with paliperidone is not recommended as

paliperidone is the active metabolite of Risperidone and the combination of the two may lead to additive active antipsychotic fraction exposure.
 Adverse Effect
 The following are all the ADRs that were reported in clinical trials and post marketing. The following terms and frequencies are applied: Very common (≥ 1/10); common (≥ 1/100); nery rare (< 1/1000), not known (cannot be estimated from the available data).
 Cardiac disorders: Tachycardia (common); atrioventricular block, bundle branch block, atrial fibrillation, sinus bradycardia, palpitations (uncommoo, ≥ 1/10); rare (≥ 1/1000; not known (cannot be estimated from the available data).
 Blood and lymphatic system disorders: Anemia, thrombocytopenia (uncommon); atrial fibrillation, sinus bradycardia, palpitations (uncommo).
 Hlood and lymphatic system disorders: Anemia, thrombocytopenia (uncommon); argunulocytopenia (rare); garanulocytosis (not known).
 Nervous system disorders: Parkinsonism (salivary hypersecretion, musculoskeletal stiffness, drooling, bradykinesia, hypokinesia, muscle rigidity), headache (very common); akathisa (restlessness, hyperkinesia, and restless leg syndrome), diszliness, speech disorder, Neuroleptic Malignant, balance disorder, tardive dyskinesia, eye disorders: Burred vision (common); conjunctivitis, ocular hyperemia, eye discharge, eye swelling, dry eye, increased lacrimation, photophobia (uncommon); reduced visual acuity, glaucoma (tree).
 Ear and ladyrinth disorders: Ear pain, tinnitus (uncommon), anasea, adominal pain, experianty, thoracic and mediastinal disorders: Dyspiea, pistaxis, cough, congestion, pharyge-laryingeal pain (common); wheezing, preunonia appration, rales, dysphonia (uncommon); dividers disorder: furgies (carno).
 Mespiratory, thoracic and mediastinal disorders: Dyspiea, pistaxis, cough, congestion, pharyge-laryingeal pain (common); whezering, preunonia apain, experiment, eye monthestiel dividers:

Skin and subcutaneous tissue disorders: Rash, erythema (common); angioedema, skin lesion, prurius, skin discoloration, alopecia, seborfnei dermatitis, dry skin, hyperkeratosis (uncommon).
 Musculoskeletal and connective tissue disorders: Arthralgia, back pain, pain in extremity (common); muscular weakness, myalgia, neck pain, joint swelling, joint stiffness, musculoskeletal chest pain (uncommon); rhabdomyolysis (rare).
 Endocrine disorders: Inappropriate anti-diurctic hormone secretion (rare).
 Metabolism and nutrition disorders: Increased appetite, decreased appetite (common); rhabdemyolysis (rare).
 Infections and infestations: Pneumonia, bronchitis, upper respiratory tract infection (crame).
 Infections and infestations: Pneumonia, bronchitis, upper respiratory tract infection.
 Vascular disorders: Hypersensitivity (uncommon); drug hypersensitivity (rare); anaphylactic reaction (nor known).
 Hepatobiliary disorders: Hypersensitivity (uncommon); drug hypersensitivity (rare); anaphylactic reaction (not known).
 Hepatobiliary disorders: Hypersensitivity (uncommon); dusfunction, erectile dysfunction, ejaculation disorders; Amenorrhea, sexual dysfunction, erectile dysfunction, ejaculation disorder, galactorrhea, gynecomastia, menstrual disorder, vaginal discharge (uncommon).
 Psychiatric disorders: Insomnia (very common); anxiety, agitation, sleep disorder (common); envination, adders, mania, decreased libido, listless, nervousness (uncommon).

(common); confusional state, mani (uncommon). DOSAGE AND ADMINISTRATION Schizophrenia

(common): conflusional state, mania, decreased notico, insuress, nervousness (uncommon).
 DOSAGE AND ADMINISTRATION
 Schizophrenia
 - Adults: Rischotic® OD may be given once daily or twice daily.
 Patients should start with 2 mg/day Rischotic® OD. The dosage may be increased on the second day to 4 mg. Most patients will benefit from daily dokes between 4 and 6 mg.
 Doses above 10 mg/day have not demonstrated superior efficacy to lower doses and may cause increased incidence of extrapyramidal symptoms. Safety of doses above 16 mg/day have not demonstrated superior efficacy to lower doses and may cause increased incidence of extrapyramidal symptoms. Safety of doses above 16 mg/day has not been evaluated, and are therefore not recommended.
 - Elderly: A starting dose of 0.5 mg twice daily ins recommended.
 - Biderly: A starting dose of 0.5 mg twice daily ins recommended.
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 - Biderly: A starting obsender of the one once daily schedule, starting with 2 mg Rischotic® OD Doses adjustmentins, if indicated, should occur at intervals of not less than 24 hours and in dosage increments of 1 mg per day. Rischotic® OD have not been investigated in patients with main cpisodes.
 - Elderly: A starting dose of 0.5 mg twice daily is recommended. This dosage can be individually adjusted with 0.5 mg twice daily is recommended. The 30 compatients being a starting with a starting with 24 mg resistent aggression in patients with moderate to severe Alzheimer's dementia.
 - A starting dose of 0.25 mg twice daily for most patients. Some patients, Nome patients,

daily. Concern may extent from 0.22 ing once daily while others may require 0.75 mg once daily. Rischotic® OD is not recommended in children less than 5 years of age, as there is no experience in children less than 5 years of age with this disorder. Renal and hepatic impairment Patients with renal impairment have less ability to eliminate the active antipsychotic fraction and have increases in plasma concentration of the free fraction of Rischotic® OD.

traction and have increases in plasma concentration of the free fraction of Kischolic[®] D. D. The spectrum of the indication, starting and consecutive dosing should be halved, and dose tirration should be slower for patients with renal or hepatic impairment. Method of administration is the starting and consecutive dosing should be halved, and dose tirration should be slower for patients with renal or hepatic impairment. Method of administration is the table of the shortprion of Rischotic[®] OD. Place the tablet on the tongue. The tablet will begin disintegrating within seconds. Water may be used if desired, No attempt should be made to divide the tablet. Upon discontinuation, gradual withdrawal is advised. Actue tablet. Upon discontinuation of high doses of antipsychotic medicines. Recurrence of psychotic synptoms may also occur, and the singence of involuntary movement disorders (such switching patients from dept antipsychotics, initiated is recommended. Also, if medically appropriate, when switching patients from dept antipsychotics, initiated Rischotic[®] OD therapy is initiated is recommended. Also, if medically appropriate, when switching patients from dept antipsychotics, initiated Rischotic[®] OD therapy in place of the next scheduled injection. The need for continuing existing anti-Parkinson medicines **CVERDOSAEE** In general, the reported signs and symptoms may include drowsiness and sedution, accurate and hepoteness.

OVERDOSAGE In general, the reported signs and symptoms may include drowsiness and sedation, tachycardia and hypotension, extrapyramidal symptoms, QT-prolongation, convulsions, Torsade de Pointes.

Iorsade de Pointes. Gastric lavage (after intubation, if the patient is unconscious) and administration of activated charcoal together with a laxative should be considered only when drug intake was less than one hour before. Cardiovascular monitoring should commence immediate-ly and should include continuous electrocardiographic monitoring to detect possible

arrhythmias. STORAGE CONDITIONS

Store below 30°C. Keep in original pack in intact conditions.

Date of revision: April 2019

This is a medicament - A medicament is a product which affects your health, and its consumption contrary to instructions is dangerous for you - Follow strictly the doctor's prescription, the method of use, and the instructions of the pharmacist who sold the medicament is the pharmacist who sold the medicament - Do not by yourself interrupt the period of treatment prescribed for you - Do not repeat the same prescription without consulting your doctor - Medicament: keep out of reach of children - Consel of Arab Health Mainsters

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