HALOPERIDOL

BOXED WARNING SECTION

WARNING ncreased Mortality in Elderly Patients with Dementia-Related Psychosis

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DESCRIPTION SECTION

Haloperidol is the first of the butyrophenone series of major tranquilizers. The chemical designation is 4-[4-(p-chlorophenyl)-4-bydroxypiperidim]-4-fluorobutyrophenone. It has the following structural formula

$$\text{CI} \underbrace{\hspace{1cm} \bigcup_{C_{21} \text{H}_{22} \text{CIPNO}_2}^{\text{HO}} \text{N} - \text{CH}_2 \text{CH}_2 \text{CH}_2}_{\text{STS-87}} \underbrace{\hspace{1cm} \bigcup_{C_{21} \text{H}_{22} \text{CIPNO}_2}^{\text{CIPNO}_2} \text{CH}_2 \text{C$$

Each balogeridol tables, USP intended for oral administration contain balogeridol, USP 5 mg or 10 mg or 20 mg, Inaddition each tables contain the following incretive ingredience; calciumstorane, dhaloc calciumplosphase dubdes, providence (PM 4.00), soulmasters by place and starch. 5 mg, D & C Leale and To A C Blee # 1 Alterianal Labe; 20 mg; FD & C Vellow #6 Alterianal Labe and D & C Red 227 Alterianal Labe.

CLINICAL PHARMACOLOGY SECTION

INDICATIONS & USAGE SECTION

NDICATIONS & USAGE SECTION

Hadoprefield is indicated for use in the management of mail festations of psychotic disorders.
Hadoprefield is indicated for the count of tics and vocal interactive. Of Tourier's Disorder in clithical
and adults, Haloprefield is indicated for the count of tics and vocal interactive of policy management
of the property of the count of th

CONTRAINDICATIONS SECTION

Haloperidol is contraindicated in severe toxic central nervous system depression or comatose states from any cause and in individuals who are hypersensitive to this drug or have Parkinson's disease.

WARNINGS SECTION

WARNINGS SECTION
Increased Mortallity in Elderly Patients with Dementia-Related Psychosis
Elderly patients with dementia-re-lated psychosis treated with antipsychotic drugs are at an increase
risk of death. Halioperidol is not approved for the treatment of patients with dementia-related psych
(see BOXED WARNING).

. Cardiovascular Effects

Cardiovascular Effects
Gases of section (I) "prolongation, and Torondes de Pointen hort born reported in patients
Gases of section (II) "I prolongation, and Torondes de Pointen hort born reported in patients
and patients (III) and the prolongation (III) and III) and prolongation (III) and III) and I

Tardwo Dyshiessa.

A syndrous constitute of potentially investrable, into hearty, dyshieric reversees my develop in higher and the electric special potential of the electric special potential pote

Whether aritysychoic drug products differ in their potential to case undive systients is unknown. Both the risk of developing under obsystema and the likelihood that six will become irreversible are sometimed to be pattern in the control of the

course of the syndrom is unknown.

Grown these consideration, antipsychotic drugs should be prescribed in a matter that is most likely to Grown these consideration, antipsychotic drugs should be prescribed in a matter that is most likely to reserved for patients who suffer from a chontic likess that, 1)1s known to respond to antipsychotic reserved to patients who suffer from a chontic likess that, 1)1s known to respond to antipsychotic likes, and, 2) for whomal-arriane, equally freshrichen known to respond to antipsychotic likes, and, 2) for whomal-arriane, equally freshrichen known to be a bread for a chost of the chost

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Neuroleptic Malignant Syndrome (NMS)

Neurolepic Maliguant Syndroms (NMS)
A potentially fault asymptom complex consortiems referred to as Neurolepic Maliguant Syndroms (NMS)
has been reported in association with antipsychoic drags. Clinical muniferations of NMS are
hyperpress, macric region, altered mean than (faciliting cannot region) and evidence of of
shripsyndrams). Additional signs may include elevanted creating phospholations, proglophismis
(thinkbomyohysis) and care rend failure.

The diagnostic evaluation of patients with this syndroms is complicated. In arriving a data genotic, it is
presented as the program of t

It a palient requires antipsychoic drug treatment after recovery from NMS, the potential reintroduction of drug therapy should be carefully considered. The patient should be carefully monitored, since recurrences of NMS have been reported. He patient should be carefully monitored, since recurrences of NMS have been reported. Hypertyrexia and heat strole, not associated with the above symptom complex, have also been reported with haloperiod.

Usage In Pregnancy Non-teratogenic Effects

Neonates exposed to antipsychotic drugs, during the flird trimester of pregnancy are at risk for exargyramidal and/or windrawal symptoms following delivery. There have been reports of agitation, hyperonia, hypotonia, terrant, roamic-terr, exprisarry obserses and feeding doctore in these enomates. These complications have varied in severity, while in some case symptoms have been self-initined, in other cases mounted have required intensive case unsupport and prolonged basisphaltandon.

Haloperidol should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Haloperiols should be used during pregames only if the potential benefit patients the potential risks in fer ferm.

Redeem given 2 is 20 itseus the usual maximum human dose of haloperiold by ord or parentered rounts for the present of the present

A number of cases of bronchopneumonia, some fatal, have followed the use of antipsycho including haloperidol. It has been postulated that lethargy and decreased semaion of this certail inhibition may lead to dehylutation, hemoconceration and reduced pulmonary vent Therefore, if the above signs and symptoms appear, especially in the elderity, the physicia intuitiate remedial therapy promphy.

Although not reported with haloperidol, decreased serum cholesterol and/or cutaneous and ocular changes have been reported in patients receiving chemically-related drugs.

Haloperidol may impair the mental and/or physical abilities required for the performance of hazardous tasks such as operating machinery or driving a motor vehicle. The ambulatory patient should be warned accordingly.

accorningity. The use of alcohol with this drug should be avoided due to possible additive effects and by

PRECAUTIONS SECTION

Lecluspeas, Neurospeata and Agranulocytosis
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In clinical sind and postumele ring requireme, events nel leukoperative moyerda have here reported
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[recluding fail case leak) has do been reported.
Possible risk factors for leukoperative moperati a relative precessing low white blood cell count (WBC)
Possible risk factors for leukoperative moperative moperative reported in the story of dreg induced leukoperative moperative relative precessing low WEG or a history
of dreg induced leukoperative moperative relative processing low WEG or a history
of dreg induced leukoperative moperative relative processing levels. Use a first single of a decline in WEG in the absence of other cannotive factors, posterior hadees USe and
frest single of a decline in WEG in the absence of other cannotive factors, posterior hadees USe and
frest single of a decline in WEG in the absence of other cannotive factors, posterior single of
infection and reason from WEG in the absence of other cannotive factors proposed in the USE and have their WBC
URapperation handle be endirativent cannotively to patients:

• with revere cardiovascular disorders, because of 8e possibility of transfert hypotemion and/or
precipitation of again plants. Should hypotemion occur and a conspecsion for required, episparities
should not be used since hadsperiods in my block its woopressor activity and paradoxical further
should not such a pressure may occur. Intend. Intend. Intend. Processing anticonvolutant medications, with a history of sciences, or with EEG absurmabilities,

- because haloperidol may lower the convulsive threshold. If indicated, adequate articonvulsant therapy should be concomitantly minimized.

 with known alleging, or with a history of allergic resections to drugs.

 receiving anticoagulates, since an isolated instance of interference occurred with the effects of one articoagulate (pleintidone).

If concominant antiparkinson medication is required, it may have to be continued after haloperidol discontinued because of the difference in excretion rates. If both are discontinued similaterously examparantial symptom may occur. The physician should keep in mind the possible increase in imaculear pressure when anticholinergic drugs, including antiparkinson agents, are administered concominantly with haloperidol.

concentrately with haloperedel. As who there are government and the langeredel may be capable of potentiating CNS depresses such as amendments, epides, and alcohol.

CNS depresses such as amendments, epides, and alcohol. In the same of the capable of potentiating the capable of the capable

When haloperidol is used to control mania in cyclic disorders, there may be a rapid mood swing to depression.

Severe neurotoxicity (rigidity, inability to walk or talk) may occur in patients with thyrotoxicosis who are also receiving antiosychotic medication, including haloneridol.

segression. Severe meutonicity (rigidory, adultity) to vall or adult my occur in patient with thyrotoxicosis who Severe meutonicity (rigidory, adultity) to vall or adult my occur in patient with thyrotoxicosis who Severe meutonicity (rigidory, adultity) to the control of the patient of the

Pregnancy: Non-teratogenic Effects

Non-transports Effects
Nonmare response to autispychotic chups, during the first trimester of pregnancy are at risk for exargy randial andro windrawal symptoms following delivery. There have been respons of agitation hypertonia, hyponian term, commiscire, respiratory distress and freeling disorder in these resonances. These complications have varied inserverity, while in some cases symptoms have been self-timated, in other cases resonance may be a complete and the self-timated in other cases resonance may require limitarity are an usin support and principle Insupstitutions. Hadoperited should be used during pregnancy only if the potential benefit is justifies the potential risks to the feats.

the fetus.

Safety and effectiveness is pediatric patients have not been established.

Gertairic Use

Safety and effectiveness in pediatric patients have not been established.

Gertairic Use

Clinical andes of hologoridol did not include sufficient numbers of subjects aged 65 and over

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ADVERSE REACTIONS SECTION

ADVAINS REACTIONS SECTION

Conflowardual Effect
Techyorida, Spoptension, and hypertension have been reported, QT prolongation and/or ventricular
arthylmstan have also been reported, in addition to ECC pattern changes compatible with the
analythmists have also deservations are reported as the process of th

psychotic patient when they go untered or when they are neared with other attripystotic drugs.

CNS HECCE

EXTLANYMANDAL SYMPTOMS (1959)— FSP during the administration of haloperiods have been

EXTLANYMANDAL SYMPTOMS (1954)— They during the administration of haloperiods have been

Parkinon-like symptom, administ, or symptomic inference in the particular of the particula

WITHDRAWAL EMERCENT NUEBOLOGICAL SIGNS formelly, patient reviewing dushers three growthers are problems with abruy discontinuation antipsychotic drugs. However, tome patients on minimum trustment experience remained policies grins after abrugs withdrawal. In certain of best cases, the dyshined movements are indistinguishable from the synthome described below under "TARDIVE DYSKINISAI" except for duration, it is not town whether gradual withdrawal of antipsychotic drugs will reduce her aim of occurrence withdrawal energent neurological signs but until further evidence becomes available, it seem eranounder up gradually withdraw are of thoughperidis.

TARDIVE DYSKINESIA

TARDIVE DYSINISIA

As with all analysochusic agents, bulgoritish has been associated with persistent dysikinesias. Turkive dysikinesia, sayadome consisting of potentially intervenible, involutary, dysikinetic nonvenent, may agene in some patients on nolsy-erine deeps or my occur after due plercy has been do continued. The risk agenes is be grower in otherly interior on big-bloose therapy, especially fermide. The risk agenes is be grower in other type and the property of the

It has been reported that fine vermicular movement of the tongue may be an early sign of tardive dyskinesia and if the medication is stopped at that time, the full syndrome may not develop. dysbatesis and if the medications is stopped at that time, the bull syndrome may not develop. TARDIVE DYSTONIAIA

Tardive dystonia, not associated with the above syndrome, has also been reported. Tardive dyston

Characterized by delayed oneset of Choreic or dystonic movement, is often persistent, and has the

OPHER CNS EFFECTS

Body as a Whole

Neuroleptic miligrant syndrome (NMS), hyperpyrexia and heat stroke have been reported with haloperidol (see WARNINGS for further information concerning NMS).

Hematologic Effects

Reminiongic Lifects

Reports have appeared citing the occurrence of mild and usually transient leukopenia and leukocytosis, minimal decreases in red blood cell counts, amenta, or a tendency toward lymphomonocytosis.
Agramalocytosis has rarely been reported to have occurred with the use of haloperidol, and then only in association with other medication.

association with other medication.

Liver Effects limpaired liver function and/or justified by the been reported.

Demandoig Recordions

Macdopopular and accretionn skin reaction and toolated cases of photosemitivity and loss of hair.

Endocrico Discontion.

Endocrine Disconfers

Lactation, breast engargement, mustalgia, menarual irregularities, gynecomusia, impotence, increa
libido, hyperglycenia, hypoghycenia and hyponatemia.

Garatinessian Effect, constigation, diarrhea, hypersall varion, dyspepsia, nussea and vomiting.

Automatic Reaction

Dry mouth, historier vision, unitary retention, diaphoresis and prinpism.

Dry mount, nutreed vision, urinary retermion, diapnoresis and priap Respiratory Effects Laryngospasm, bronchospasm and increased depth of respiration. Special Senses

Cataracts, retinopathy and visual disturbances.
Postmarketing Events

Hyperammonemia has been reported in a 5½ year old child with citrullinemia, an inherited disorder of ammonia excretion, following treatment with haloperidol.

Maniferations in ingernal, the symptoms of overdestage would be an estaggeration of inover pharmacologic effects and anderest reactions, the most primaries of which would be: 1) severe estaggerated reactions, 2) by physical may have found the severe enemyle produce a shack like learn. The excrappy readiled reaction would be marries by mancular weakens or rigidity and a generalized or localized enterer as estaggerated to the control of th

Transmer

Gantic Issage or induction of emesis should be carried out immediately followed by administration activated claimed. Since there is no specific author, became its principle superview. A plant as its concess, by taches onsity. Recipitatry of perspectives on proceedings of the process of the concess of the process of the proce

is normal. Severe arrhythmias should be treated with appropriate anti-arrhythmic measures.

DOSAGE & ADMINISTRATION SECTION

DOSAGE A DMINISTRATION SECTION

There is considered a cuitation imagine to a patient in the amount of medication required for treatment. As with all antipoychoic charge, do sage should be individualized according to the receive are response of each patient. Dosage adjustmens, either upward on downward, should be carried out as rapidly as practicable to achieve optimum therepeate control.

To determine the initial dosage, consideration should be given to the patient's age, severity of illness, previous response to other antipoychoic drugs, and any concominant medication or disease state. Challens, debilitude of graitary patients, we will as those with a latency of adverse reactions to include a control of the control

.........Austinistration INITIAL DOSAGE RANGE Adults

Moderate Symptomatology	0.5 mg to 2 mg b.i.d. or t.i.d.
Severe Symptomatology	3 mg to 5 mg b.i.d. or t.i.d.

To achieve prompt control, higher doses may be required in some cases.

Geriatric or Debilitated Patients
Chronic or Resistant Patients
Patients who remain assertly disturbed or implementally controlled 0.5 mg to 2 mg b.i.d. or t.i.d. 3 mg to 5 mg b.i.d. or t.i.d.

Children
The following recommendation apply to children between the ages of 3 and 12 years (weight range 15 to 40 kg). Haloporidol is not invended for children under 3 years old. Therapy should begin at the lowest does possible (4.5 mg per day.) If required, the does should be increased by an increment of 0.5 mg as 6 to 7 dy inventive under the description of the descript

Psychotic Disorders

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Maintenance Dosage

Upon achieving a satisfactory therapeutic response, dosage should then be gradually reduced lowest effective maintenance level.

Switchover Procedure

Switchners Procedure

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The earl foundated pupping the injectable as soon as practicable, in the debrere of biocordiability of the early of

HOW SUPPLIED SECTION

Haloperidol Tablets USP, 5 mg are green, capsule-shaped, flat-faced, beveled-edge tablets debossed with the logo of 'ZC', '07' and partial bisect, on one side and plain on the other side and are supplied as follows:

NDC 68382-079-10 in bottles of 1000 tablets
Haloperidol Tablets USP, 10 mg are light green, capsule-shaped, flas-faced, beveled-edge tablets debossed with the logo of 2CC, '08' and partial bisect, on one side and plain on the other side and are supplied as follows:

sebusced with the logic of ZCC, '00' and jurital bisest, on one side and plain on the other side and are supplied as follows:

NDC 6832-080-06 in houties of 30 tables.

NDC 6832-080-06 in houties of 100 tables.

NDC 6832-080-06 in houties of 1000 tables.

NDC 6832-080-06 in houties of 2000 tables.

NDC 6832-080-06 in houties of 30 tables.

NDC 6832-080-06 in houties of 30 tables.

NDC 6832-080-06 in houties of 30 tables.

NDC 6832-080-06 (70' to 77'9') list best USP Controlled Room Temperature).

Dispense in sight, light-resistant container.

All trademates are the property of 7-plus group.

Call your decur for medical advice about side effects. You may report side effects to FDA at 1-800-1744-0408.

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL





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