### Haloneridol Tablets, USP

### WARNING

Increased Mortally in Electry Parkens with Dementia-Related Psychosis. Electry patients with Generation-standar pytics in transit with astroychotic drugs are at an increased risk of death. Analyses of seventeen placebo- controlled trials (modal duration of 10 weeks), largely in patients stating applied analyses of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a paying a large part of the sevent as a large part of the sevent as a large part of the sevent which the first goal of the sevent was a large part of the sevent with the first goal of the sevent was the sevent with the first first goal of the part of the sevent with the first first goal of the part of the part of the sevent with the first first goal of the sevent with the first forther of part of the sevent with the first first goal of the sevent with the first goal of the sevent with the first goal of the sevent with the first goal of the sevent goal of the sevent with the first goal of the sevent goal of the sevent with the first goal of the sevent goal of the s

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

Each haloperidol tablet, USP intended for oral administration contain haloperidol, USP 5 mg or 10 mg or 20 mg, in addition each tablet contains the following interive ingredients: calcium testurate, dibasic calcium phosphase dibayletae, providee (PVF NS), sodimustavelly speciale and states, Ns mg Do R C mg C new C ne

### CLINICAL PHARMACOLOGY

has not been clearly established.

### INDICATIONS AND USAGE

Haloperidol is indicated for use in the management of manifestations of psychotic disorders Hadopericol is indicated for use in the management of numletastions of psychotic disorders. Hadopericol is indicated for the course of its cand wood unteracts of Tourette's Disorder in children and adults. Hadopericol is effective for the renament of severe behavior problems in children of contabler, explored hypersecticality (which cannot be accusated for by immediate provisoration), emitted the provisoration of the contabler explored for the provisoration of the contabler explored in the provisoration of the contabler explored in the contable explored in the con

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.

Increased Mortality in Elderly Patients with Dementia-Related Psychosis
Elderly patients with dementia-related psychosis treated with antipsychotic drugs are at an increase of each. Haloperidol is not approved for the treatment of patients with dementia-related psyckes BOXED WARNING.

### Cardiovas cular Effects

Cardiavascular Ellects
Cases of suded neith, (T-ypolong ation, and Torsades de Pointes have been reported in patients receiving halpeerisch. Higher han recommende dosses of any formulation of halpeeriola Japane as be associated with a labyer risk of (T-yeolongation and Torsades de Pointes. Although cases have been reported even in the absence of presisposing factors, particular canton is advised in retenting patients with other (T-yeolongia conditions (inclined jetective) in thistance [particularly plosplations and hypomagnes-mish], drugs Insonso prolong (T, underlying cardiac absorrantities, hypothyroidton, and traillail long (T-yeofatowa).

### Tardive Dyskinesia

A syndrom consisting of potentially irreversible, involuntary, dyskinetic movements may develop in patients treated with autipsychotic drugs. Although the prevalence of the syndrome appears to be highest among the deferly, especially lederly women, it is impossible to rely upon prevalence estimates to predict, as the inception of autipsychotic treatment, which patients are likely to develop the syndrome. Whether antipsychotic drugs produces differ in their potential coanse turked welysteria is untravow. whether anapychotic rung products duriet in meri potentia in coanse tarture obstaness is unanown. Both the risk of evel poing tarfet opsianess in and the likelihood that it will become irreversible are believed to increase as the duration of treatment and the total cumulative dose of artipsychotic drug; administered to the patient increase. However, the syndrome can develop, although much less commonly, after relatively brief treatment periods at low doses.

commons, after returney brief reasures periods all tow doses.

There is no known reasured for established cases of surfive depkinesia, although the syndrome may result, partially or completely, if antipsychotic treatment is withdrawn. Antipsychotic treatment is withdrawn. Antipsychotic treatment, itself, however, may suppress (operatiful suppress) the signs and syndroms of the syndroms and thereby may possibly must the underlying process. The effect that symptomatic suppression has upon the long-term course of the syndroms is ulmanous.

course of the syndrome is unknown. Given these condicions, arisys, both drugs should be prescribed in a numer that is most likely to minimize the occurrence of surdive dyskiresia. Chronic antipoychoic treatment should generally be reserved for patients who suffer from a chronic illens shat, it) is known to respond a surject hook drugs, and, 2) for whom alternative, equally effective, has potentially less harmful restateness are not vanished or appropriate, in patients who of regrite chronic treatment, the smillest does and the shortest duration of reatment producing a satisfactory clinical response should be sought. The need for continued examents should be reasonable proficiously.

If signs and symptoms of turdive dyskirensia appear in a againet on antipsychotic, drug discontinuation should be considered. However, some guidents may require treatment despite the presence of the should be considered. However, some guidents may require the entired they like the presence of the control of

## Neuroleptic Malignant Syndrome (NMS)

A potentially faal symptom complex sometimes referred to as Neurolegic Malignant Symfrom (NMS) has been reported in association with antipsychoic dungs. Clinical manifestations of NMS are more proposed in association with antipsychoic dungs. Clinical manifestations of NMS are more proposed in a substance of the substance of the

(thabdomydysis) and acut rental failure. The diagnostic evaluation of patients with this syndrome is complicated. In arriving at a diagnosis, it is important to identify cases where the clinical presentation includes both serious medical illness (e.g., appenunous, systemic reference), are increased or inadequality tened exarasymmatida signs and symptom (EFS). Other important considerations in the differential diagnosis include certain symptom (EFS) to the important considerations in the magnetic diagnosis include certain symptomic results of the consideration of the symptomic consideration of the consideration of the symptomic design and solver drugs not essential to concurrent therapy, 2) intends design outsides for essential consistency and other drugs not essential to concurrent therapy. 2) intends the discontinuities of supplying the consistency of the

If a patient requires, artipsychoic 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.

Hyperpressix and best stroke, not associated with the above symptom complex, have also been reported with hadoperiod.

## Falls

Fault Haloperidol may cause somrolence, postural hypotension, motor and sensory instability, which may lead to falls and, consequently, fractures or other injuries. For patients with diseases, conditions, or medications that could exacerbate these effects, complete fall risk assessments when initiating antipsychotic treatment and recurrently for patients on long-term antipsychotic therapy.

# Usage In Pregnancy Non-teratogenic Effects

Non-teraingeme Eltects

Nonates expose on antipsychotic drugs, during the third trimester of pregnancy are at risk for exrapyramidal and/or withdrawal symptoms following delivery. There have been reports of agitation, hypertonia, hypotrala, termor, sommle-ter, espriaturely distress and feeding distored in these reconstant. These complications have varied in severity, while in some cases symptoms have been self-limited, in other cases rouses have required intensive care unit support and prolonged hospitalization.

ridol should be used during pregnancy only if the potential benefit justifies the potential risk to the feats.

Rodens given 2 to 20 times the usual maximum human dose of haloperidol by oral or parenteral routes showed an increase in incidence of resorption, reduced fertility, delayed delivery and pay mortality. No habe the control of the contr

is no evidence to relate this phenomenon to predictable human risk for most of these agents. There are no well consolied studies with haloppiral to in preguato women. There are reports, however, of cases of limb multiormations observed following material use of haloperiold along with other drugs which have suspected entangenic potential during the list interior of pregnanty. Canal relationships were not established in these cases. Size such experience does not exclude the possibility of feat during due to haloppirals, this days should be used during pregnary or in women light to be tour pregnant only if the beautific clearly justifies a potential risk to the feats. Indians should not be moved during drug resultses.

## Combined Use of Haloperidol and Lithium

Combined Use of Halperfield and Lithium
An excephalipation cythrone (characterised by weakers, lethurg), fewr, trendomness and
confusion, extragorantial symptomic, brainey wiss, lethurg), fewr, trendomness and
confusion, extragorantial symptomic, brainey wiss, lethurg, fewr and extra express, BUN, and FBS) followed
by irreverable be rain damage has occurred in a few patients reason with hillium plus halperfield. A
cansal relationship between these events and the conconstant administration of lithium and halperfield
has not been established, however, patients receiving such confidencings should be munitored
closely for early evidence of neurological toxicity and resument discontinued promptly if such signs
appear.

supear.

General

A muther of cases of bronchopperumoria, some fatal, have followed the use of antipsychoic drugs
including hadoperishel. It has been postulated that behaving and decreased sensation of thirst due to
central inhibition may lead to desiydation, hemoconcentation and reduced pulmonary ventilation.

Therefore, if the above signs and symposts appear, especially in the elderity, the physician should
institute remedial therapy promptly.

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

\*\*Halometed have missing the means algority or bruscle abilities required for the performance of hazard

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.

### PRECAUTIONS

### Leukopenia, Neutropenia and Agranulocytos is

In clinical trial and postmarketing experience, events of leukopenia/neutropenia have been reported temporally related to antipsychotic agents, including haloperidol tablets USP. Agramulocytosis (including faatla cases) has also been reported.

Passible risk factors for feelingensiateuropens irritade previoting low white blood cell count ONLO.

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Passible risk factors for feelingensiateuropens and previous feelingens and the count of the county of the passible risk feelingensiateuropens also blood the cell county of the passible feelingens and the county of the passible respective for the passible respective for the passible feelingensiateuropens and the carefully monitored for feet or other symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly if such symptoms or signs of infection and restored promptly in such symptoms.

- Integrated to that the Cuttery, and the Cuttery and the Cutter
- should be used.
  receiving anticomodusat medications, with a history of seizanes, or with EEG abnormalities,
  because haloperidol imp lower the convolvive threshold. If indicated, adequate anticonvolusat
  with bown allergies, or with a history of allergic reactions to drugs.
  receiving anticogalizate, since an isolated instance of interference occurred with the effects of one
  anticogalizat (hearthfulore).

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 simultaneously, excreting the continued of the physician should been just find the possible increase in exemption of the physician should be printed the possible increase in concominantly with haloperidol.

As with other antipsychotic agents, it should be noted that haloperidol may be capable of potentiating CNS depressants such as anesthetics, opiates, and alcohol.

In a study of 12 schänophereit patients condunistatered haloperidol and rifampia, plasmu haloperidol levels were decreased by a mun oil 70% and mean scores on the first Psychaint; Edung Scale were severed to the study of the s

.

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

depression.

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

Severe neuronacci y riguiny, analimi yo waxe for also pin occur in paneres wun in pronauccious wan born mategieria portuda el follogierialo von fonta in he Anes-Salmonelli neirconoma di sevicioni assay, Negative or inconsistera positive findings, hore bero hostined in in vitro and in vivo studies of effects of habiperido en chromosome structure and maher. The availables opportugies evicines is considered too inconsistera be conclusive at this time. Carcinogeneity sudues using oral haboperidol were conducted in Wistar rats (dooed at up to 5 mg/kg daily for 24 months) and in Alians Swiss ratse (dosed at up to 5 mg/kg daily for 24 months) and in Alians Swiss ratse (dosed at up to 5 mg/kg daily for 24 months) and in Alians Swiss ratse (dosed at up to 5 mg/kg daily for 18 months), in the rat study in high dose male and female groups, these animals of did not have a greater incidence of tumors study in high-dose male and female groups, these animals of did not have a greater incidence of tumors hand cornoral animals. Therefore, although not portunal, this study because great incidence of tumors hand to the control of the cont

sans many, wase surer was a summanally significant increase in patients; gland resplication, limited rifect, no statistically significant differences in incredences of total names or specific insurers to repetit the production of the composition of the end update of the production of the production of the end update of the production of the production of the end update of the end update of the production of the end update of the end update

Non-teruspiern L. [Icts]
Nonates exposed to antipsychotic drugs, during the third trimester of pregnancy are at risk for extrapyramidal and/or withdrawal symptoms following delivery. There have been reports of agitat hypertonia, hypotonia, terror, seromice, respiratory distress and feeding disorder in these some These complications have varied in severity, while in some cases symptoms have been self-limite other cases roomsten shave required insertsive care until support and prolonged thospitalization.

Pediatric Use Safety and effec

### Geriatric Use

Geranic USE

(Clinical studies of haloperidol did not include sufficient numbers of subjects aged 65 and over to
determize whether they respond differently from younger subjects. Other reported clinical experience
has not consistently disentified differences in responses between the delively any younger patients.
However, the prevalence of tardite dyslatensia appears to be highest among the elderly, especially
elderly yourne (see WANINOS: Tardite Dyslatensia), also he plasmacolimics of haloperidol in
gertaint; quietes generally warrans the use of lower dises (see DOSAGE AND
AMMINISTRATIOS).

## ADVERSE REACTIONS

## Cardiovas cular Effects

Cardiavascular Effects
Tackycardia, hypotenion, and hypertenion have been reported. QT prolongation audior ventricular arrhydrains have also been reported, in addition to ECC pattern changes compatible with the oppolymorphous configuration of terstaced espoints, and may occur more frequently with high doses and in predisposed patterns (see WARNINGS and PRECAUTIONS).

Classes of sudden and unexpected deals have been reported in a sociation with the administration of an haloperitod. The nature of the evidence makes it impossible to desertaine definitively what role, if any haloperitod played the no account of the reported cases. The postability that haloperitod cased deals caused under the control of the reported cases. The postability that haloperitod cased deals cross to the control of the reported cases. The postability that haloperitod cased deals reported cases. The postability that haloperitod cased deals reported cases. The postability that haloperitod cased deals on the control of the reported deals and user period deals may occur in postability that haloperitod changed deals are controlled to the control of the postability that haloperitod cased deals on the control of the reported deals and user period deals may occur in problement of the postability that haloperitod change deals are controlled to the control of the postability that haloperitod change deals are controlled to the contro

PS) stone, present CNS IIIerds

EXTEAPYRAMIDAL SYMPTOMS (EPS) — EPS during the administration of haloperiod have been reported frequently, often during the first few days of reament. EPS can be categorized generally as Parkisson-like symptom, shadnish, or dystonia furchaling opisthotonous and conlogyric crisis). While all can occur at relatively low doses, they occur more frequently and wing greater severity in higher dones. The symptoms may be common frequently and one of the stone of the ston

Dynama. Class effect: Symptoms of dystoria, prolonged abnormal contractions of muscle groups, may occur in succeptible individuals during the first few days of restment. Dystoric symptoms include: spaces of the succeptible individuals during the first few days of restment. Dystoric symptoms include: spaces of the present process of the succeptible of the succeptible of the succeptible of the succeptible of the present process of the succeptible of the succeptible of the succeptible of the process of the succeptible of suc

## WITHDRAWAL EMERGENT NEUROLOGICAL SIGNS

Graphily, patient receiving shorterm therapy experience reproduces with abuse discontinual configuration of the production of the producti

## TARDIVE DYSKINESIA

LARLIVE DYSMINESIA

As with all antiportotic agents, haloperidol has been associated with persistent dyskinesias. Tardir dyskinesia, a syndrome consisting of potentially irreversible, involutatory, obslancic movemens, may appear intone patients on long-term fleeny or may occur after drug theory has been discontinued, and appear intone patients on long-term fleeny or may occur after drug theory has been discontinued, by the contractive of the cont

movement of extremnes and the trust.

There is no known effective treatment for tardive dyskinesia; antiparkinton agents usually do not adleviate the symptom of this symbon. It is unggreated that all antipsychotic agents be discontinued and approximately the symbol of the symbol of the symbol of the agent, or which no additinest antipsychotic agent, shi syndrome mybe maked.

It has been reported that first 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 my not develop.

## TARDIVE DYSTONIA

Tardive dystoria, not associated with the above syndrome, has also been reported. Tardive dystoriance the delayed onset of choreic or dystoric movemens, is often persistent, and has the potential of becoming irreversible.

## OTHER CNS EFFECTS

Incomnia, residensenses, anxiety, euphoria, agliation, drowsiness, depression, lethargy, headache, confusion, verigo, grand mil seizures, exacerbation of psychotic symptoms including halluc inations and catantic-libe behavioral states which may be responsive to drug withdrawal and/or treatment with anticholinergic drugs.

## Body as a Whole

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

## Hematologic Effects

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

## Liver Effects

aired liver function and/or jaundice have been reported.

Maculopapular and acneiform skin reactions and isolated cases of photosensitivity and loss of hair.

Lactation, breast engorgement, mastalgia, menstrual irregularities, gynecomastia, impotence, increased libido, hyperglycemia, hypoglycemia and hyponatremia.

### Gas trointes tinal Effects

Anorexia, constination, diarrhea, hypersalivation, dyspensia, nausea and vomiting,

Res piratory 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.

## OVERDOSAGE

OVERIONAGE

Munifications
In general, the symptoms of overdosage would be an exaggeration of known pharmacologic effects and
adverse reactions, the most prominer of which would be: 1) severe extrapyratidel reactions, 2)
hypotension, or 3) seldation. The painest would appear commotor with respiratory depression and
the promiser of the principle of the painest would appear commotor with respiratory depression and
cuttom could be marifestly by muccular vealuses or rigidity and a generalized or localized remote and
demonstrated by the adventised and against types respectively. With accidental overdosage, hypertension
are than hypotension cocurred in a two-year old child. The risk of ECC changes accided with
refer to ADVERSE REACTIONS).

refer to ADVERSE REACTIONS).

Trenument

Gamic longe or induction of enersis should be carried on immediately followed by administration of servined charged to thought on the property of the

### DOSAGE AND ADMINISTRATION

DOSAGE AND ADMINISTRATION
There is condicated variation from patient to patient in the amount of medication required for treatment. As with all antipsychotic drugs, dosage should be individualized according to the needs and treatment. As with all antipsychotic drugs, dosage should be individualized according to the needs and engaging and approximate the content of the needs and required area. To determine the initial dosage, consideration should be given to the patient; sage, severity of illuses. To determine the initial dosage, consideration should be given to the patient; sage, severity of illuses. Children, debilized or geriatric patients, as well as those with a history of adverse reactions to obtained with more grantal dosage adjustments and at lower dosage levels, as recommended below. Clinical experience suggests the following recommendations:

Oral Administration 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 severely disc 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 recommendations apply to children between the ages of 3 and 12 years (weight range 15 or 04 kg). Histoperidol is not intended for children under 3 years old. Therapy should begin at the lowest done possible (0.5 mg per day). It required, the doses should be increased by an increment of 0.5 mg at 5 or 7 day intended, under the quark of the distinction of the distincti

The total dose may be divided, to be given b.i.d. or t.i.d.

Psychotic Disorders

1.05 mg/hg/day to 0.15 mg/hg/day

1.05 mg/hg/day

1.05 mg/hg/day to 0.15 mg/hg/day

1.05 mg

Maintenance Dosage

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

Invest effective munteauxe rever.

The oral formshould supplant he injectable as soon as practicable. In the absence of binovailability studies establishing blooquivalence between these two dosage forms, the following guidelines for dosage are suggested. For an initial approximation of the total daily dose required, the parenteral dose administered in the preceding 24 hours may be used. Since this dose in only a minial estimate, it is sentiance for the preceding 24 hours may be used. Since this dose in only a minial estimate, it is sentiance of such as a sential proposal position of the preceding and adverse effects, be carried on periodically for the first several days following the initiation of switchers, this two, dose, adaptament, either quant of ordowned, can be quickly accomplished. Depending on the patient's clinical stums, the first oral dose should be given within 12-24 hours following the last parenteral dose.

Haloperidol Tables 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 63832-079-01 in bottles of 100 tables

NDC 63832-079-10 in bottles of 1000 tables

NDC 63832-079-10 in bottles of 1000 tables

Haloperiola Tables USP, 100 gar et light gerne, aguale-shaped, flas faced, beweled-edge tables, debosed with the logo of 272.00 and partial bisect, on one side and plain on the other side and are supplied as follows.

NDC 63832-080-0 in bottles of 30 tables

NDC 63832-080-10 in bottles of 1000 tables

NDC 63832-080-10 in bottles of 1000 tables

NDC 63832-080-10 in bottles of 1000 tables

Haloperidol Tablets USP, 20 mg are coral, capsule-shaped, flat-faced, beveled-edge tablets debossed with the logo of 'ZC', '09' and bisect on one side and plain on the other side and are supplied as follows:

NDC 68382-081-06 in bottles of 30 tablets NDC 68382-081-01 in bottles of 100 tablets

NDC 68382-081-01 in butles of 100 tables

Size at 30° to 20° (60° to 77°) [See USP Controlled RoomTemperature].

Dispense may be 20° to 20° to

Manufactured by: Cadila Healthcare Ltd.

Ahmedabad, India

Distributed by: Zvdus Pharmaceuticals USA Inc.

nnington, NJ 08534 Pennington, NJ 06534 Rev.: 11/16 Revision Date : 2016/11/12



haloperidol tablet							
Product Informati	on						
Product Type	HUMAN PRESCRIPTION DRUG Rem Code (Se				e) NDC:57046-3	29(ND	C:68382-075
Route of Administrati	on	ORAL.					
Active Ingredient/	Active Moie	ty					
Ingredient Name  HALOPERIDOL (UNI: 16292F813D) (BALOPERIDOL - UNI: 16292F813D)					Basis of Strength Streng MALOPERDOL 5 mg		Strength 5 mg
Inactive Ingredien	ts						
		Ingredient Name					Strength
CALCIUM STEARATE (							
D&C YELLOW NO. 10							
		DRATE (UNI: O7TSZ97GEP)					
FD&C BLUE NO. 1-AL PO VIDONE K30 (UNIE		(UNR J9EQA352JM)				_	
		A POTATO (UNE 5856/3G2)				-	
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ALUMINUM O XIDE (UP		)				_	
Product Character	istics						
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						10 mm	
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Labeler - Contract Pharmacy Services-PA (945429777) 
 Establishment
 Name
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 Business Operations

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 Revised: 6/2020
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